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HELICOBACTER PYLORI PROTEINS USEFUL FOR VACCINES AND DIAGNOSTICS

Abstract:

Abstract of WO9318150

Helicobacter pylori is known to cause or be a cofactor in type B gastritis, peptic ulcers, and gastric tumors. In both developed and developing countries, a high percentage of people are infected with this bacterium. The present invention relates generally to certain H. pylori proteins, to the genes which express these proteins, and to the use of these proteins for diagnostic and vaccine applications. Specifically, molecular cloning, nucleotide, and amino acid sequences for the H. pylori cytotoxin (CT), the "Cytotoxin Associated Immunodominant" (CAI) antigen, and the heat shock protein (hsp60) are described herein. Data supplied from the esp@cenet database - Worldwide

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(71) Applicant (for all designated States except US): BIOCINE SCLAVO S.P.A. [IT/IT]; Via Fiorentina, 1, I-53100 Siena (IT).

(72) Inventors; and

(75) Inventors/Applicants (for US only): COVACCI, Antonello [IT/IT]: BUGNOLI, Massimo [IT/IT]: Via Fiorentina, 1, I-53100 Siena (IT). TELFORD, John [GB/IT]; Via Fiorentina, 1, I-43100 Siena (IT). MACCHIA, Giovanni [IT/IT]; RAPPUOLI, Rino [IT/IT]; Via Fiorentina, 1, I-53100 Siena (IT).

(74) Agent: HALLYBONE, Huw, George; Carpmaels & Ransford, 43 Bloomsbury Square, London WC1A 2RA (GB).

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Helicobacter pylori is known to cause or be a cofactor in type B gastritis, peptic ulcers, and gastric tumors. In both developed and developing countries, a high percentage of people are infected with this bacterium. The present invention relates generally to certain H. pylori proteins, to the genes which express these proteins, and to the use of these proteins for diagnostic and vaccine applications. Specifically, molecular cloning, nucleotide, and amino acid sequences for the H. pylori cytotoxin (CT), the "Cytotoxin Associated Immunodominant" (CAI) antigen, and the heat shock protein (hsp60) are described herein.

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HELICOBACTER PYLORI PROTEINS USEFUL FOR VACCINES AND DIAGNOSTICS

BACKGROUND OF THE INVENTION

1. Field of the Disclosure

The present invention relates generally to certain Helicobacter pylori proteins, to the genes which express these proteins, and to the use of these proteins for diagnostic and vaccine applications.

2. Brief Description of Related Art

Helicobacter pylori is a curved, microaerophilic, gram negative bacterium that has been isolated for the first 10 time in 1982 from stomach biopsies of patients with chronic gastritis, Warren et al., Lancet i:1273-75 Originally Campylobacter pylori, it has been named recognized to be part of a separate genus 15 Helicobacter, Goodwin et al., Int. J. Syst. Bacteriol. 39:397-405 (1989). The bacterium colonizes the human gastric mucosa, and infection can persist for decades. During the last few years, the presence of the bacterium has been associated with chronic gastritis type B, a condition that may remain asymptomatic in most infected persons but 20 increases considerably the risk of peptic ulcer and gastric adenocarcinoma. The most recent studies strongly suggest that <u>H. pylori</u> infection may be either a cause or a cofactor of type B gastritis, peptic ulcers, and gastric tumors, see e.g., Blaser, Gastroenterology 93:371-83 (1987); Dooley et 25 al., New Engl. J. Med. 321:1562-66 (1989); Parsonnet et al., New Engl. J. Med. 325:1127-31 (1991). H. pylori is believed to be transmitted by the oral route, Thomas et al., Lancet i:340, 1194 (1992), and the risk of infection increases with age, Graham et al., Gastroenterology 30 100:1495-1501 (1991), and is facilitated by crowding, Drumm et al., New Engl. J. Med. 4322:359-63 (1990); Blaser, Clin. Infect. Dis. 15:386-93 (1992). In developed countries, the presence of antibodies against <u>H. pylori</u> antigens increases from less than 20% to over 50% in people 30 and 60 years old 35 respectively, Jones et al., Med. Microbio. 22:57-62 (1986); Morris et al., N.Z. Med. J. 99:657-59 (1986), while in

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developing countries over 80% of the population are already infected by the age of 20, Graham et al., Digestive Diseases and Sciences 36:1084-88 (1991).

The nature and the role of the virulence factors of <u>H. pylori</u> are still poorly understood. The factors that have been identified so far include the flagella that are probably necessary to move across the mucus layer, see e.g., Leying et al., Mol. Microbiol. 6:2863-74 (1992); the urease that is necessary to neutralize the acidic environment of the stomach and to allow initial colonization, see e.g., Cussac et al., J. Bacteriol. 174:2466-73 (1992); Perez-Perez et al., J. Infect. Immun. 60:3658-3663 (1992); Austin et al., J. Bacteriol. 174:7470-73 (1992); PCT Publ. No. WO 90/04030; and a high molecular weight cytotoxic protein formed by monomers allegedly having a molecular weight of 87 kDa that causes formation of vacuoles in eukaryotic epithelial cells and is produced by H. pylori strains associated with disease, see e.g., Cover et al., J. Bio. Chem. 267:10570-75 (1992) (referencing a "vacuolating toxin" with a specified 23 amino acid N-terminal sequence); Cover et al., J. Clin. Invest. 90:913-18 (1992); Leunk, Rev. Infect. Dis. 13:5686-89 (1991). Additionally, the following is also known.

H. pylori culture supernatants have been shown by different authors to contain an antigen with a molecular 25 weight of 120, 128, or 130 kDa, Apel et al., Aentralblat fur Bakteriol. Microb. und Hygiene 268:271-76 (1988); Crabtree et al., J. Clin. Pathol 45:733-34 (1992); Cover et al., Infect. Immun. 58:603-10 (1990); Figura et al., H. pylori, 30 gastritis and peptic ulcer (eds. Malfrtheiner et al.), Springer Verlag, Berlin (1990). Whether the difference in size of the antigen described was due to interlaboratory differences in estimating the molecular weight of the same protein, to the size variability of the same antigen, or to actual different proteins was not clear. No nucleotide or 35 amino acid sequence information was given about the protein. This protein is very immunogenic in infected humans because specific antibodies are detected in sera of virtually all patients infected with H. pylori, Gerstenecker et al., Eur.

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J. Clin. Microbiol. 11:595-601 (1992).

H. pylori heat shock proteins (hsp) have been described, Evans et al., Infect. Immun. 60:2125-27 (1992) (44 amino acid N-terminal sequence and a molecular weight of about 62 kDa); Dunn et al., Infect. Immun. 60:1946-51 (1992) (33 amino acids found in the N-terminal sequence and a molecular weight of about 54 kDa); Austin et al., J. Bacteriol. 174:7470-73 (1992) (37 amino acids found in the N-terminal sequence and a molecular weight of about 60 kDa). Austin et al. suggest that these are, in fact, the same protein with identical amino acid sequences at their N-terminus.

For examples of diagnostic tests based on <u>H. pylori</u> lysates or semipurified antigens, see Evans et al., Gastroenterology 96:1004-08 (1989); U.S. 4,882,271; PCT Publ. No. WO 89/08843 (all relating to compositions and assays containing the same having high molecular weight antigens (300-700 kDa) from the outer membrane surface with urease activity); EPO Publ. No. 329 570 (relating to antigenic compositions for detecting <u>H. pylori</u> antibodies having fragments of at least one fragment from the group 63, 57, 45, and 31 kDa).

The percentage of people infected by <u>H. pylori</u>, either in a symptomatic or an asymptomatic form, is very high in both developing and developed countries, and the cost of hospitalization and therapy makes desirable the development of both <u>H. pylori</u> vaccines and further diagnostic tests for this disease.

30 . SUMMARY OF THE INVENTION

The present invention describes nucleotide and amino acid sequences for three major <u>H. pylori</u> proteins. Specifically, these are the cytotoxin, the "Cytotoxin Associated Immunodominant" (CAI) antigen, and the heat shock protein. None of the complete amino acid sequences for these proteins has been known, nor have their genes been identified. The present invention pertains to not only these purified proteins and their genes, but also recombinant materials associated therewith, such as vectors

WO 93/18150 PCT/EP93/00472

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and host cells. The understanding at the molecular level of the nature and the role of these proteins and the availability of recombinant production has important implications for the development of new diagnostics for <u>H. pylori</u> and for the design of vaccines that may prevent <u>H. pylori</u> infection and treat disease.

As such, these proteins can be used in both vaccine and diagnostic applications. The present invention includes methods for treating and diagnosing those diseases associated with <u>H. pylori</u>. As <u>H. pylori</u> has been associated with type B gastritis, peptic ulcers, and gastric adenocarcinoma, it is hoped that the present invention will assist in early detection and alleviation of these disease states. Currently, diagnosis relies mostly on endoscopy and histological staining of biopsies; existing immunoassays are based on <u>H. pylori</u> lysates or semi-purified antigens. Given the heterogeneity found in such assays, correlation with disease state is not yet well established. Thus, the potential for recombinant antigen-based immunoassays, as well as nucleic acid assays for disease detection, is great. At present, there is no commercial vaccine for H. pylori infection or treatment. A recombinant vaccine is thus an object of the present invention.

25 BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is the nucleotide sequence for the cytotoxin (CT) protein.

Fig. 2 is the amino acid sequence for the cytotoxin (CT) protein.

Fig. 3 is a map of the <u>cai</u> gene for the CAI protein and summary of the clones used to identify and sequence this gene.

Fig. 4 is the nucleotide and amino acid sequences of the CAI antigen.

Fig. 5 is the nucleotide and amino acid sequences of the heat shock protein (hsp).

DETAILED DESCRIPTION OF THE INVENTION

A. General Methodology

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The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. techniques are explained fully in the literature. See e.g., Sambrook, et al., MOLECULAR CLONING; A LABORATORY MANUAL, SECOND EDITION (1989); DNA CLONING, VOLUMES I AND II (D.N Glover ed. 1985); OLIGONUCLEOTIDE SYNTHESIS (M.J. Gait ed, 1984); NUCLEIC ACID HYBRIDIZATION (B.D. Hames & S.J. Higgins eds. 1984); TRANSCRIPTION AND TRANSLATION (B.D. Hames & S.J. Higgins eds. 1984); ANIMAL CELL CULTURE (R.I. Freshney ed. 1986); IMMOBILIZED CELLS AND ENZYMES (IRL Press, 1986); B. Perbal, A PRACTICAL GUIDE TO MOLECULAR CLONING (1984); the series, METHODS IN ENZYMOLOGY (Academic Press, Inc.); GENE TRANSFER VECTORS FOR MAMMALIAN CELLS (J.H. Miller and M.P. Calos eds. 1987, Cold Spring Harbor Laboratory), Methods in Enzymology Vol. 154 and Vol. 155 (Wu and Grossman, and Wu, eds., respectively), Mayer and Walker, eds. (1987), IMMUNOCHEMICAL METHODS IN CELL AND MOLECULAR BIOLOGY (Academic London), Press, Scopes, (1987),PROTEIN PURIFICATION: PRINCIPLES AND PRACTICE, Second Edition (Springer-Verlag, N.Y.), and HANDBOOK OF EXPERIMENTAL IM-MUNOLOGY, VOLUMES I-IV (D.M. Weir and C. C. Blackwell eds 1986).

Standard abbreviations for nucleotides and amino acids are used in this specification. All publications, patents, and patent applications cited herein are incorporated by reference.

30 <u>B. Definitions</u>

"Cytotoxin" or "toxin" of <u>H. pylori</u> refers to the protein, and fragments thereof, whose nucleotide sequence and amino acid sequences are shown in Figs. 1 and 2, respectively, and their derivatives, and whose molecular weight is about 140 kDa. This protein serves as a precursor to a protein having an approximate weight of 100 kDa and having cytoxic activity. The cytotoxin causes vacuolation and death of a number of eukaryotic cell types and has been purified from <u>H. pylori</u> culture supernatants. Additionally,

the cytotoxin is proteinaceous and has an apparent molecular mass determined by gel filtration of approximately 950-972 kDa. Denaturing gel electrophoresis of purified material previously revealed that the principal component of the 950-972 kDa molecule was allegedly a polypeptide of apparent molecular mass of 87 kDa, Cover et al., J. Biol. Chem. 267:10570-75 1992). It is suggested herein, however, that the previously described 87 kDa results from either the further processing of the 100 kDa protein or from proteolytic degradation of a larger protein during purification.

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The "Cytotoxin Associated Immunodominant" (CAI) antigen refers to that protein, and fragments thereof, whose amino acid sequence is described in Fig. 4 and derivatives thereof. This is an hydrophilic, surface-exposed protein having a molecular weight of approximately 120-132 kDa, preferably 128-130 kDa, produced by clinical isolates. The size of the gene and of the encoded protein varies in different strains by a mechanism that involves duplication of regions internal to the gene. The clinical isolates that do not produce the CAI antigen, do not have the cai gene, and are also unable to produce an active cytotoxin. association between the presence of the cai gene and cytotoxicity suggests that the product of the cai gene is necessary for the transcription, folding, export or function of the cytotoxin. Alternatively, both the cytotoxin (CT) and the cai gene are absent in noncytotoxic strains. This would imply some physical linkage between the two genes. A peculiar property of the CAI antigen is the size variability, suggesting that the cai gene is continuously changing. The CAI antigen appears to be associated to the cell surface. This suggests that the release of the antigen in the supernatant may be due to the action of proteases present in the serum that may cleave either the antigen itself, or the complexes that hold the CAI antigen associated to the bacterial surface. Similar processing activities may release the antigen during in vivo growth. The absence of a typical leader peptide sequence suggests the presence of an independent export system.

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"Heat shock protein" (hsp) refers to the <u>H. pylori</u> protein, and fragments thereof, whose amino acid sequence is given in Fig. 5 and derivatives thereof, and whose molecular weight is in the range of 54-62 kDa, preferably about 58-60 kDa. This hsp belongs to the group of Gram negative bacteria heat shock proteins, hsp60. In general, hsp are among the most conserved proteins in all living organisms, either prokaryotic and eukaroytic, animals and plants, and the conservation is spread along the whole sequence. This high conservation suggests a participation of the whole sequence at the functional structure of the protein that can be hardly modified without impairing its activity.

Examples of proteins that can be used in the present invention include polypeptides with minor amino acid variations from the natural amino acid sequence of the protein; in particular, conservative amino acid replacements are contemplated. Conservative replacements are those that take place within a family of amino acids that are related in their side chains. Genetically encoded amino acids are generally divided into four families: (1) acidic = aspartate, glutamate; (2) basic = lysine, arginine, histidine; (3) non-polar = alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan; and (4) uncharged polar = glycine, asparagine, glutamine, cystine, serine, threonine, tyrosine. Phenylalanine, tryptophan, and tyrosine are sometimes classified jointly as aromatic amino acids. For example, it is reasonably predictable that an isolated replacement of a leucine with an isoleucine or valine, an aspartate with a glutamate, a threonine with a serine, or a similar conservative replacement of an amino acid with a structurally related amino acid will not have a major effect on the biological activity. Polypeptide molecules having substantially the same amino acid sequence as the protein but possessing minor amino acid substitutions that do not substantially affect the functional aspects are within the definition of the protein.

A significant advantage of producing the protein by recombinant DNA techniques rather than by isolating and

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purifying a protein from natural sources is that equivalent quantities of the protein can be produced by using less starting material than would be required for isolating the protein from a natural source. Producing the protein by recombinant techniques also permits the protein to be isolated in the absence of some molecules normally present in cells. Indeed, protein compositions entirely free of any trace of human protein contaminants can readily be produced because the only human protein produced by the recombinant non-human host is the recombinant protein at issue. Potential viral agents from natural sources and viral components pathogenic to humans are also avoided.

The term "recombinant polynucleotide" as used herein intends a polynucleotide of genomic, cDNA, semisynthetic, or synthetic origin which, by virtue of its origin or manipulation: (1) is not associated with all or a portion of a polynucleotide with which it is associated in nature, (2) is linked to a polynucleotide other than that to which it is linked in nature, or (3) does not occur in nature. Thus, this term also encompasses the situation wherein the <u>H. pylori</u> bacterium genome is genetically modified (e.g., through mutagenesis) to produce one or more altered polypeptides.

The term "polynucleotide" as used herein refers to a polymeric form of a nucleotide of any length, preferably deoxyribonucleotides, and is used interchangeably herein with the terms "oligonucleotide" and "oligomer." The term refers only to the primary structure of the molecule. Thus, this term includes double- and single-stranded DNA, as well as antisense polynucleotides. It also includes known types of modifications, for example, the presence of labels which are known in the art, methylation, end "caps," substitution of one or more of the naturally occurring nucleotides with an analog, internucleotide modifications such as, for example, replacement with certain types of uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoamidates, carbamates, etc.) or charged linkages phosphorothioates, phosphorodithioates, etc.), introduction of pendant moieties, such as, for example,

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proteins (including nucleases, toxins, antibodies, signal peptides, poly-L-lysine, etc.), intercalators (e.g., acridine, psoralen, etc.), chelators (e.g., metals, radioactive species, boron, oxidative moieties, etc.), alkylators (e.g., alpha anomeric nucleic acids, etc.).

By "genomic" is meant a collection or library of DNA molecules which are derived from restriction fragments that have been cloned in vectors. This may include all or part of the genetic material of an organism.

By "cDNA" is meant a complimentary mRNA sequence that hybridizes to a complimentary strand of mRNA.

As used herein, the term "oligomer" refers to both primers and probes and is used interchangeably herein with the term "polynucleotide." The term oligomer does not connote the size of the molecule. However, typically oligomers are no greater than 1000 nucleotides, more typically are no greater than 500 nucleotides, even more typically are no greater than 250 nucleotides; they may be no greater than 100 nucleotides, and may be no greater than 75 nucleotides, and also may be no greater than 50 nucleotides in length.

The term "primer" as used herein refers to an oligomer which is capable of acting as a point of initiation of synthesis of a polynucleotide strand when used under appropriate conditions. The primer will be completely or substantially complementary to a region of the polynucleotide strand to be copied. Thus, under conditions conducive to hybridization, the primer will anneal to the complementary region of the analyte strand. Upon addition of suitable reactants, (e.g., a polymerase, nucleotide triphosphates, and the like), the primer will be extended by the polymerizing agent to form a copy of the analyte strand. The primer may be single-stranded or alternatively may be partially or fully double-stranded.

The terms "analyte polynucleotide" and "analyte strand" refer to a single- or double-stranded nucleic acid molecule which is suspected of containing a target sequence, and which may be present in a biological sample.

As used herein, the term "probe" refers to a structure comprised of a polynucleotide which forms a hybrid

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structure with a target sequence, due to complementarily of at least one sequence in the probe with a sequence in the target region. The polynucleotide regions of probes may be composed of DNA, and/or RNA, and/or synthetic nucleotide analogs. Included within probes are "capture probes" and "label probes".

As used herein, the term "target region" refers to a region of the nucleic acid which is to be amplified and/or detected. The term "target sequence" refers to a sequence with which a probe or primer will form a stable hybrid under desired conditions.

The term "capture probe" as used herein refers to a polynucleotide probe comprised of a single-stranded polynucleotide coupled to a binding partner. The single-stranded polynucleotide is comprised of a targeting polynucleotide sequence, which is complementary to a target sequence in a target region to be detected in the analyte polynucleotide. This complementary region is of sufficient length and complementarily to the target sequence to afford a duplex of stability which is sufficient to immobilize the analyte polynucleotide to a solid surface (via the binding partners). The binding partner is specific for a second binding partner; the second binding partner can be bound to the surface of a solid support, or may be linked indirectly via other structures or binding partners to a solid support.

The term "targeting polynucleotide sequence" as used herein refers to a polynucleotide sequence which is comprised of nucleotides which are complementary to a target nucleotide sequence; the sequence is of sufficient length and complementarily with the target sequence to form a duplex which has sufficient stability for the purpose intended.

The term "binding partner" as used herein refers to a molecule capable of binding a ligand molecule with high specificity, as for example an antigen and an antibody specific therefor. In general, the specific binding partners must bind with sufficient affinity to immobilize the analyte copy/complementary strand duplex (in the case of

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capture probes) under the isolation conditions. Specific binding partners are known in the art, and include, for example, biotin and avidin or streptavidin, IgG and protein numerous Α, the known receptor-ligand couples, complementary polynucleotide strands. In the case of complementary polynucleotide binding partners, the partners are normally at least about 15 bases in length, and may be at least 40 bases in length; in addition, they have a content of Gs and Cs of at least about 40% and as much as about 60%. The polynucleotides may be composed of DNA, RNA, or synthetic nucleotide analogs.

The term "coupled" as used herein refers to attachment by covalent bonds or by strong non-covalent interactions (e.g., hydrophobic interactions, hydrogen bonds, etc.). Covalent bonds may be, for example, ester, ether, phosphoester, amide, peptide, imide, carbon-sulfur bonds, carbon-phosphorus bonds, and the like.

The term "support" refers to any solid or semi-solid surface to which a desired binding partner may be anchored. Suitable supports include glass, plastic, metal, polymer gels, and the like, and may take the form of beads, wells, dipsticks, membranes, and the like.

The term "label" as used herein refers to any atom or moiety which can be used to provide a detectable (preferably quantifiable) signal, and which can be attached to a polynucleotide or polypeptide.

As used herein, the term "label probe" refers to a polynucleotide probe which is comprised of a targeting polynucleotide sequence which is complementary to a target sequence to be detected in the analyte polynucleotide. This complementary region is of sufficient length and complementarily to the target sequence to afford a duplex comprised of the "label probe" and the "target sequence" to be detected by the label. The label probe is coupled to a label either directly, or indirectly via a set of ligand molecules with high specificity for each other, including multimers.

The term "multimer," as used herein, refers to linear or branched polymers of the same repeating

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single-stranded polynucleotide unit different or single-stranded polynucleotide units. At least one of the units has a sequence, length, and composition that permits it to hybridize specifically to a first single-stranded nucleotide sequence of interest, typically an analyte or a polynucleotide probe (e.g., a label probe) bound to an In order to achieve such specificity and analyte. stability, this unit will normally be at least about 15 nucleotides in length, typically no more than about 50 nucleotides in length, and preferably about 30 nucleotides in length; moreover, the content of Gs and Cs will normally be at least about 40%, and at most about 60%. In addition to such unit(s), the multimer includes a multiplicity of units that are capable of hybridizing specifically and stably to a second single-stranded nucleotide of interest, typically a labeled polynucleotide or another multimer. These units are generally about the same size and composition as the multimers discussed above. When a multimer is designed to be hybridized to another multimer, the first and second oligonucleotide units are heterogeneous (different), and do not hybridize with each other under the conditions of the selected assay. Thus, multimers may be label probes, or may be ligands which couple the label to the probe.

A "replicon" is any genetic element, e.g., a plasmid, a chromosome, a virus, a cosmid, etc. that behaves as an autonomous unit of polynucleotide replication within a cell; i.e., capable of replication under its own control. This may include selectable markers.

"PCR" refers to the technique of polymerase chain reaction as described in Saiki, et al., Nature 324:163 (1986); and Scharf et al., Science (1986) 233:1076-1078; and U.S. 4,683,195; and U.S. 4,683,202.

As used herein, x is "heterologous" with respect to y if x is not naturally associated with y in the identical manner; i.e., x is not associated with y in nature or x is not associated with y in the same manner as is found in nature.

"Homology" refers to the degree of similarity

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between x and y. The correspondence between the sequence from one form to another can be determined by techniques known in the art. For example, they can be determined by a direct comparison of the sequence information of the polynucleotide. Alternatively, homology can be determined by hybridization of the polynucleotides under conditions which form stable duplexes between homologous regions (for example, those which would be used prior to S₁ digestion), followed by digestion with single-stranded specific nuclease(s), followed by size determination of the digested fragments.

A "vector" is a replicon in which another polynucleotide segment is attached, so as to bring about the replication and/or expression of the attached segment.

"Control sequence" refers to polynucleotide sequences which are necessary to effect the expression of coding sequences to which they are ligated. The nature of such control sequences differs depending upon the host organism; in prokaryotes, such control sequences generally include promoter, ribosomal binding site, and transcription termination sequence; in eukaryotes, generally, such control sequences include promoters and transcription termination sequence. The term "control sequences" is intended to include, at a minimum, all components whose presence is necessary for expression, and may also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences.

"Operably linked" refers to a juxtaposition wherein the components so described are in a relationship permitting them to function in their intended manner. A control sequence "operably linked" to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequences.

An "open reading frame" (ORF) is a region of a polynucleotide sequence which encodes a polypeptide; this region may represent a portion of a coding sequence or a total coding sequence.

A "coding sequence" is a polynucleotide sequence

WO 93/18150

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PCT/EP93/00472

which is translated into a polypeptide, usually via mRNA, when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a translation start codon at the 5'-terminus and a translation stop codon at the 3'-terminus. A coding sequence can include, but is not limited to, cDNA, and recombinant polynucleotide sequences.

As used herein, the term "polypeptide" refers to a polymer of amino acids and does not refer to a specific length of the product; thus, peptides, oligopeptides, and proteins are included within the definition of polypeptide. This term also does not refer to or exclude post expression modifications of polypeptide, the for example, glycosylations, acetylations, phosphorylations and the like. Included within the definition are, for polypeptides containing one or more analogs of an amino acid (including, for example, unnatural amino acids, etc.), polypeptides with substituted linkages, as well as other modifications known in the art, both naturally occurring and non-naturally occurring.

A polypeptide or amino acid sequence "derived from" a designated nucleic acid sequence refers to a polypeptide having an amino acid sequence identical to that of a polypeptide encoded in the sequence, or a portion thereof wherein the portion consists of at least 3-5 amino acids, and more preferably at least 8-10 amino acids, and even more preferably at least 11-15 amino acids, or which is immunologically identifiable with a polypeptide encoded in the sequence. This terminology also includes a polypeptide expressed from a designated nucleic acid sequence.

"Immunogenic" refers to the ability of a polypeptide to cause a humoral and/or cellular immune response, whether alone or when linked to a carrier, in the presence or absence of an adjuvant. "Neutralization" refers to an immune response that blocks the infectivity, either partially or fully, of an infectious agent.

"Epitope" refers to an antigenic determinant of a peptide, polypeptide, or protein; an epitope can comprise 3 or more amino acids in a spatial conformation unique to the

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epitope. Generally, an epitope consists of at least 5 such amino acids and, more usually, consists of at least 8-10 such amino acids. Methods of determining spatial conformation of amino acids are known in the art and include, for example, x-ray crystallography and 2-dimensional nuclear magnetic resonance. Antibodies that recognize the same epitope can be identified in a simple immunoassay showing the ability of one antibody to block the binding of another antibody to a target antigen.

"Treatment," as used herein, refers to prophylaxis and/or therapy (i.e., the modulation of any disease symptoms). An "individual" indicates an animal that is susceptible to infection by <u>H. pylori</u> and includes, but is not limited to, primates, including humans. A "vaccine" is an immunogenic, or otherwise capable of eliciting protection against <u>H. pylori</u>, whether partial or complete, composition useful for treatment of an individual.

The <u>H. pylori</u> proteins may be used for producing antibodies, either monoclonal or polyclonal, specific to the proteins. The methods for producing these antibodies are known in the art.

"Recombinant host cells", "host cells," "cells," "cell cultures," and other such terms denote, for example, microorganisms, insect cells, and mammalian cells, that can be, or have been, used as recipients for recombinant vector or other transfer DNA, and include the progeny of the original cell which has been transformed. It is understood that the progeny of a single parental cell may not necessarily be completely identical in morphology or in genomic or total DNA complement as the original parent, due to natural, accidental, or deliberate mutation. Examples for mammalian host cells include Chinese hamster ovary (CHO) and monkey kidney (COS) cells.

Specifically, as used herein, "cell line," refers
to a population of cells capable of continuous or prolonged
growth and division in vitro. Often, cell lines are clonal
populations derived from a single progenitor cell. It is
further known in the art that spontaneous or induced changes
can occur in karyotype during storage or transfer of such

clonal populations. Therefore, cells derived from the cell line referred to may not be precisely identical to the ancestral cells or cultures, and the cell line referred to includes such variants. The term "cell lines" also includes immortalized cells. Preferably, cell lines include nonhybrid cell lines or hybridomas to only two cell types.

As used herein, the term "microorganism" includes prokaryotic and eukaryotic microbial species such as bacteria and fungi, the latter including yeast and filamentous fungi.

"Transformation", as used herein, refers to the insertion of an exogenous polynucleotide into a host cell, irrespective of the method used for the insertion, for example, direct uptake, transduction, f-mating or electroporation. The exogenous polynucleotide may be maintained as a non-integrated vector, for example, a plasmid, or alternatively, may be integrated into the host genome.

referring to a polypeptide or nucleotide sequence, that the indicated molecule is present in the substantial absence of other biological macromolecules of the same type. The term "purified" as used herein preferably means at least 75% by weight, more preferably at least 85% by weight, more preferably at least 85% by weight, more preferably at least 98% by weight, of biological macromolecules of the same type present (but water, buffers, and other small molecules, especially molecules having a molecular weight of less than 1000, can be present).

30 C. Nucleic Acid Assays

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Using as a basis the genome of <u>H. pylori</u>, polynucleotide probes of approximately 8 nucleotides or more can be prepared which hybridize with the positive strand(s) of the RNA or its complement, as well as to cDNAs. These polynucleotides serve as probes for the detection, isolation and/or labeling of polynucleotides which contain nucleotide sequences, and/or as primers for the transcription and/or replication of the targeted sequences. Each probe contains a targeting polynucleotide sequence, which is comprised of

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nucleotides which are complementary to a target nucleotide sequence; the sequence is of sufficient length and complementarily with the sequence to form a duplex which has sufficient stability for the purpose intended. For example, if the purpose is the isolation, via immobilization, of an analyte containing a target sequence, the probes will contain a polynucleotide region which is of sufficient length and complementarily to the targeted sequence to afford sufficient duplex stability to immobilize the analyte on a solid surface under the isolation conditions. example, also, if the polynucleotide probes are to serve as primers for the transcription and/or replication of target sequences, the probes will contain a polynucleotide region of sufficient length and complementarily to the targeted sequence to allow for replication. For example, also, if the polynucleotide probes are to be used as label probes, or are to bind to multimers, the targeting polynucleotide region would be of sufficient length and complementarily to form stable hybrid duplex structures with the label probes and/or multimers to allow detection of the duplex. probes may contain a minimum of about 4 contiguous nucleotides which are complementary to the targeted sequence; usually the oligomers will contain a minimum of about 8 continuous nucleotides which are complementary to the targeted sequence, and preferably will contain a minimum of about 14 contiguous nucleotides which are complementary to the targeted sequence.

The probes, however, need not consist only of the sequence which is complementary to the targeted sequence. They may contain additional nucleotide sequences or other moieties. For example, if the probes are to be used as primers for the amplification of sequences via PCR, they may contain sequences which, when in duplex, form restriction enzyme sites which facilitate the cloning of the amplified sequences. For example, also, if the probes are to be used as "capture probes" in hybridization assays, they will be coupled to "binding a partner" defined as Preparation of the probes is by means known in the art, including, for example, by methods which include excision,

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transcription or chemical synthesis.

D. Expression Systems

Once the appropriate <u>H. pylori</u> coding sequence is isolated, it can be expressed in a variety of different expression systems; for example those used with mammalian cells, baculoviruses, bacteria, and yeast.

i. Mammalian Systems

Mammalian expression systems are known in the art. A mammalian promoter is any DNA sequence capable of binding mammalian RNA polymerase and initiating the downstream (3') transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiating region, which is usually placed proximal to the 5' end of the coding sequence, and a TATA box, usually located 25-30 base pairs (bp) upstream of the transcription initiation site. The TATA box is thought to direct RNA polymerase II to begin RNA synthesis at the correct site. A mammalian promoter will also contain an upstream promoter element, usually located within 100 to 200 bp upstream of the TATA box. An upstream promoter element determines the rate at which transcription is initiated and can act in either orientation, Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed (1989).

Mammalian viral genes are often highly expressed and have a broad host range; therefore sequences encoding mammalian viral genes provide particularly useful promoter sequences. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter (Ad MLP), and herpes simplex virus promoter. In addition, sequences derived from non-viral genes, such as the murine metallotheionein gene, also provide useful promoter sequences. Expression may be either constitutive or regulated (inducible), depending on the promoter can be induced with glucocorticoid in hormone-responsive cells.

The presence of an enhancer element (enhancer), combined with the promoter elements described above, will usually increase expression levels. An enhancer is a regulatory DNA sequence that can stimulate transcription up to 1000-fold when linked to homologous or heterologous

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promoters, with synthesis beginning at the normal RNA start site. Enhancers are also active when they are placed upstream or downstream from the transcription initiation site, in either normal or flipped orientation, or at a distance of more than 1000 nucleotides from the promoter, Maniatis et al., Science 236:1237 (1989); Alberts et al. Molecular Biology of the Cell, 2nd ed (1989). Enhancer elements derived from viruses may be particularly useful, because they usually have a broader host range. include the SV40 early gene enhancer, Dijkema et al (1985) EMBO J. 4:761, and the enhancer/promoters derived from the long terminal repeat (LTR) of the Rous Sarcoma Virus, Gorman et al. (1982) Proc. Natl. Acad. Sci. 79:6777, and from human cytomegalovirus, Boshart et al. (1985) Cell 41:5221. Additionally, some enhancers are regulatable and become active only in the presence of an inducer, such as a hormone or metal ion, Sassone-Corsi et al. (1986) Trends Genet. 2:215; Maniatis et al. (1987) Science 236:1237.

A DNA molecule may be expressed intracellularly in mammalian cells. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by the ATG start codon. If desired, the N-terminus may be cleaved from the protein by in vitro incubation with cyanogen bromide.

Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in mammalian cells. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either in vivo or in vitro. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. The adenovirus tripartite leader is an example of a leader sequence that provides for secretion of a foreign protein in mammalian cells.

Usually, transcription termination and

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polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. The 3' terminus of the mature mRNA is formed by site-specific post-transcriptional cleavage and polyadenylation, Birnstiel et al. (1985) Cell 41:349; Proudfoot and Whitelaw (1988) "Termination and 3' end processing of eukaryotic RNA. In Transcription and splicing (ed. B.D. Hames and D.M. Glover); Proudfoot (1989) Trends Biochem. Sci. sequences direct the These 14:105. transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminator/polyadenylation signals include those derived from SV40, Sambrook et al (1989), Molecular Cloning: A Laboratory Manual.

Some genes may be expressed more efficiently when introns (also called intervening sequences) are present. Several cDNAs, however, have been efficiently expressed from vectors that lack splicing signals (also called splice donor and acceptor sites), see e.g., Gething and Sambrook (1981) Nature 293:620. Introns are intervening noncoding sequences within a coding sequence that contain splice donor and acceptor sites. They are removed by a process called "splicing," following polyadenylation of the primary transcript, Nevins (1983) Annu. Rev. Biochem. 52:441; Green (1986) Annu. Rev. Genet. 20:671; Padgett et al. (1986) Annu. Rev. Biochem. 55:1119; Krainer and Maniatis (1988) "RNA splicing," In Transcription and splicing (ed. B.D. Hames and D.M. Glover).

Usually, the above-described components, comprising a promoter, polyadenylation signal, and transcription termination sequence are put together into expression constructs. Enhancers, introns with functional splice donor and acceptor sites, and leader sequences may also be included in an expression construct, if desired. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (e.g., plasmids) capable of stable maintenance in a host, such as mammalian cells or bacteria. Mammalian replication systems include those

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derived from animal viruses, which require trans-acting factors to replicate. For example, plasmids containing the replication systems of papovaviruses, such as SV40, Gluzman (1981) Cell 23:175, or polyomavirus, replicate to extremely high copy number in the presence of the appropriate viral T antigen. Additional examples of mammalian replicons include those derived from bovine papillomavirus and Epstein-Barr virus. Additionally, the replicon may have two replication systems, thus allowing it to be maintained, for example, in mammalian cells for expression and in a procaryotic host for cloning and amplification. Examples of such mammalian-bacteria shuttle vectors include pMT2, Kaufman et al. (1989) Mol. Cell. Biol. 9:946, and pHEBO, Shimizu et al. (1986) Mol. Cell. Biol. 6:1074.

The transformation procedure used depends upon the host to be transformed. Methods for introduction of heterologous polynucleotides into mammalian cells are known in the art and include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

Mammalian cell lines available as hosts for expression are known in the art and include many immortalized cell lines available from the American Type Culture Collection (ATCC), including but not limited to, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human hepatocellular carcinoma cells (e.g., Hep G2), and a number of other cell lines.

ii. Baculovirus Systems

The polynucleotide encoding the protein can also be inserted into a suitable insect expression vector, and is operably linked to the control elements within that vector. Vector construction employs techniques which are known in the art.

Generally, the components of the expression system include a transfer vector, usually a bacterial plasmid, which contains both a fragment of the baculovirus genome,

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and a convenient restriction site for insertion of the heterologous gene or genes to be expressed; a wild type baculovirus with a sequence homologous to the baculovirus-specific fragment in the transfer vector (this allows for the homologous recombination of the heterologous gene in to the baculovirus genome); and appropriate insect host cells and growth media.

After inserting the DNA sequence encoding the protein into the transfer vector, the vector and the wild type viral genome are transfected into an insect host cell where the vector and viral genome are allowed to recombine. The packaged recombinant virus is expressed and recombinant plaques are identified and purified. Materials and methods baculovirus/insect cell expression systems for commercially available in kit form from, inter alia, Invitrogen, San Diego CA ("MaxBac" kit). These techniques are generally known to those skilled in the art and fully described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987) (hereinafter "Summers and Smith").

Prior to inserting the DNA sequence encoding the protein into the baculovirus genome, the above-described components, comprising a promoter, leader (if 'desired), coding sequence of interest, and transcription termination sequence, are usually assembled into an intermediate transplacement construct (transfer vector). This construct may contain a single gene and operably linked regulatory elements; multiple genes, each with its owned set of operably linked regulatory elements; or multiple genes, regulated by the same set of regulatory elements. Intermediate transplacement constructs are often maintained in a replicon, such as an extrachromosomal element (e.g., plasmids) capable of stable maintenance in a host, such as a bacterium. The replicon will have a replication system, thus allowing it to be maintained in a suitable host for cloning and amplification.

Currently, the most commonly used transfer vector for introducing foreign genes into AcNPV is pAc373. Many other vectors, known to those of skill in the art, have also

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been designed. These include, for example, pVL985 (which alters the polyhedrin start codon from ATG to ATT, and which introduces a BamHI cloning site 32 basepairs downstream from the ATT; see Luckow and Summers, Virology (1989) 17:31.

The plasmid usually also contains the polyhedron polyadenylation signal (Miller et al. (1988) Ann. Rev. Microbiol., 42:177) and a procaryotic ampicillin-resistance (amp) gene and origin of replication for selection and propagation in E. coli.

Baculovirus transfer vectors usually contain a baculovirus promoter. A baculovirus promoter is any DNA sequence capable of binding a baculovirus RNA polymerase and initiating the downstream (5' to 3') transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A baculovirus transfer vector may also have a second domain called an enhancer, which, if present, is usually distal to the structural gene. Expression may be either regulated or constitutive.

Structural genes, abundantly transcribed at late times in a viral infection cycle, provide particularly useful promoter sequences. Examples include sequences derived from the gene encoding the viral polyhedron protein, Friesen et al., (1986) "The Regulation of Baculovirus Gene Expression," in: The Molecular Biology of Baculoviruses (ed. Walter Doerfler); EPO Publ. Nos. 127 839 and 155 476; and the gene encoding the p10 protein, Vlak et al., (1988), J. Gen. Virol. 69:765.

DNA encoding suitable signal sequences can be derived from genes for secreted insect or baculovirus proteins, such as the baculovirus polyhedrin gene (Carbonell et al. (1988) Gene, 73:409). Alternatively, since the signals for mammalian cell posttranslational modifications (such as signal peptide cleavage, proteolytic cleavage, and phosphorylation) appear to be recognized by insect cells, and the signals required for secretion and nuclear

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accumulation also appear to be conserved between the invertebrate cells and vertebrate cells, leaders of non-insect origin, such as those derived from genes encoding human α-interferon, Maeda et al., (1985), Nature 315:592; human gastrin-releasing peptide, Lebacq-Verheyden et al., (1988), Molec. Cell. Biol. 8:3129; human IL-2, Smith et al., (1985) Proc. Nat'l Acad. Sci. USA, 82:8404; mouse IL-3, (Miyajima et al., (1987) Gene 58:273; and human glucocerebrosidase, Martin et al. (1988) DNA 7:99, can also be used to provide for secretion in insects.

A recombinant polypeptide or polyprotein may be expressed intracellularly or, if it is expressed with the proper regulatory sequences, it can be secreted. Good intracellular expression of nonfused foreign proteins usually requires heterologous genes that ideally have a short leader sequence containing suitable translation initiation signals preceding an ATG start signal. If desired, methionine at the N-terminus may be cleaved from the mature protein by in vitro incubation with cyanogen bromide.

Alternatively, recombinant polyproteins or proteins which are not naturally secreted can be secreted from the insect cell by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in insects. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the translocation of the protein into the endoplasmic reticulum.

After insertion of the DNA sequence and/or the gene encoding the expression product precursor of the protein, an insect cell host is co-transformed with the heterologous DNA of the transfer vector and the genomic DNA of wild type baculovirus — usually by co-transfection. The promoter and transcription termination sequence of the construct will usually comprise a 2-5kb section of the baculovirus genome. Methods for introducing heterologous DNA into the desired site in the baculovirus virus are known in the art. (See Summers and Smith; Ju et al. (1987); Smith

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et al., Mol. Cell. Biol. (1983) 3:2156; and Luckow and Summers (1989)). For example, the insertion can be into a gene such as the polyhedrin gene, by homologous double crossover recombination; insertion can also be into a restriction enzyme site engineered into the desired baculovirus gene. Miller et al., (1989), Bioessays 4:91.

The DNA sequence, when cloned in place of the polyhedrin gene in the expression vector, is flanked both 5' and 3' by polyhedrin-specific sequences and is positioned downstream of the polyhedrin promoter.

The newly formed baculovirus expression vector is subsequently packaged into an infectious recombinant baculovirus. Homologous recombination occurs at low frequency (between about 1% and about 5%); thus, the majority of the virus produced after cotransfection is still wild-type virus. Therefore, a method is necessary to identify recombinant viruses. An advantage of the expression system is a visual screen allowing recombinant viruses to be distinguished. The polyhedrin protein, which is produced by the native virus, is produced at very high levels in the nuclei of infected cells at late times after viral infection. Accumulated polyhedrin protein forms occlusion bodies that also contain embedded particles. These occlusion bodies, up to 15 μm in size, are highly refractile, giving them a bright shiny appearance that is readily visualized under the light microscope. Cells infected with recombinant viruses lack occlusion bodies. To distinguish recombinant virus from wild-type virus, the transfection supernatant is plaqued onto a monolayer of insect cells by techniques known to those skilled in the Namely, the plaques are screened under the light art. microscope for the presence (indicative of wild-type virus) or absence (indicative of recombinant virus) of occlusion bodies. "Current Protocols in Microbiology" Vol. 2 (Ausubel et al. eds) at 16.8 (Supp. 10, 1990); Summers and Smith; Miller et al. (1989).

Recombinant baculovirus expression vectors have been developed for infection into several insect cells. For example, recombinant baculoviruses have been developed for,

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inter alia: Aedes aegypti, Autographa californica, Bombyx mori, Drosophila melanogaster, Spodoptera frugiperda, and Trichoplusia ni (PCT Pub. No. WO 89/046699; Carbonell et al., (1985) J. Virol. 56:153; Wright (1986) Nature 321:718; Smith et al., (1983) Mol. Cell. Biol. 3:2156; and see generally, Fraser, et al. (1989) In Vitro Cell. Dev. Biol. 25:225).

Cells and cell culture media are commercially available for both direct and fusion expression of heterologous polypeptides in a baculovirus/expression system; cell culture technology is generally known to those skilled in the art. See, e.g., Summers and Smith.

The modified insect cells may then be grown in an appropriate nutrient medium, which allows for stable maintenance of the plasmid(s) present in the modified insect host. Where the expression product gene is under inducible control, the host may be grown to high density, and expression induced. Alternatively, where expression is constitutive, the product will be continuously expressed into the medium and the nutrient medium must be continuously circulated, while removing the product of interest and augmenting depleted nutrients. The product may be purified by such techniques as chromatography, e.g., HPLC, affinity chromatography, ion exchange chromatography, electrophoresis; density gradient centrifugation; solvent extraction, or the like. As appropriate, the product may be further purified, as required, so as to remove substantially any insect proteins which are also secreted in the medium or result from lysis of insect cells, so as to provide a product which is at least substantially free of host debris, e.g., proteins, lipids and polysaccharides.

In order to obtain protein expression, recombinant host cells derived from the transformants are incubated under conditions which allow expression of the recombinant protein encoding sequence. These conditions will vary, dependent upon the host cell selected. However, the conditions are readily ascertainable to those of ordinary skill in the art, based upon what is known in the art.

iii. Bacterial Systems

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Bacterial expression techniques are known in the A bacterial promoter is any DNA sequence capable of art. binding bacterial RNA polymerase and initiating the downstream (3") transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A bacterial promoter may also have a second domain called an operator, that may overlap an adjacent RNA polymerase binding site at which RNA synthesis begins. The operator permits negative regulated (inducible) transcription, as a gene repressor protein may bind the operator and thereby inhibit transcription of a specific gene. Constitutive expression may occur in the absence of negative regulatory elements, such as the operator. In addition, positive regulation may be achieved by a gene activator protein binding sequence, which, if present is usually proximal (5') to the RNA polymerase binding sequence. An example of a gene activator protein is the catabolite activator protein (CAP), which helps initiate transcription of the lac operon in E. coli, Raibaud et al. (1984) Annu. Rev. Genet. 18:173. Regulated expression may therefore be either positive or negative, thereby either enhancing reducing or transcription.

Sequences encoding metabolic pathway enzymes provide particularly useful promoter sequences. Examples include promoter sequences derived from sugar metabolizing enzymes, such as galactose, lactose (lac), Chang et al. (1977) Nature 198:1056, and maltose. Additional examples include promoter sequences derived from biosynthetic enzymes such as tryptophan (trp), Goeddel et al. (1980) Nuc. Acids Res. 8:4057; Yelverton et al. (1981) Nucl. Acids Res. 9:731; U.S. 4,738,921; EPO Publ. Nos. 036 776 and 121 775. The glaotamase (bla) promoter system, Weissmann (1981) "The cloning of interferon and other mistakes." In Interferon 3 (ed. I. Gresser), bacteriophage lambda PL, Shimatake et al. (1981) Nature 292:128, and T5, U.S. 4,689,406, promoter

systems also provide useful promoter sequences.

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In addition, synthetic promoters which do not occur in nature also function as bacterial promoters. For example, transcription activation sequences of one bacterial or bacteriophage promoter may be joined with the operon sequences of another bacterial or bacteriophage promoter, creating a synthetic hybrid promoter, U.S. 4,551,433. For example, the tac promoter is a hybrid trp-lac promoter comprised of both trp promoter and lac operon sequences that is regulated by the <u>lac</u> repressor, Amann et al. (1983) Gene 25:167; de Boer et al. (1983) Proc. Natl. Acad. Sci. 80:21. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. A naturally occurring promoter of nonbacterial origin can also be coupled with a compatible RNA polymerase to produce high levels of expression of some genes in prokaryotes. bacteriophage The RNA polymerase/promoter system is an example of a coupled promoter system, Studier et al. (1986) J. Mol. Biol. 189:113; Tabor et al. (1985) Proc Natl. Acad. Sci. 82:1074. In addition, a hybrid promoter can also be comprised of a bacteriophage promoter and an E. coli operator region (EPO Publ. No. 267 851).

In addition to a functioning promoter sequence, an efficient ribosome binding site is also useful for the expression of foreign genes in prokaryotes. In E. coli, the ribosome binding site is called the Shine-Dalgarno (SD) sequence and includes an initiation codon (ATG) and a sequence 3-9 nucleotides in length located 3-11 nucleotides upstream of the initiation codon, Shine et al. (1975) Nature 254:34. The SD sequence is thought to promote binding of mRNA to the ribosome by the pairing of bases between the SD sequence and the 3' and of E. coli 16S rRNA, Steitz et al. "Genetic signals and nucleotide sequences in (1979)messenger RNA." In <u>Biological Regulation and Development:</u> Gene Expression (ed. R.F. Goldberger). To express eukaryotic genes and prokaryotic genes with weak ribosomebinding site, Sambrook et al. (1989), Molecular Cloning: A

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Laboratory Manual.

A DNA molecule may be expressed intracellularly. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus will always be a methionine, which is encoded by the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by in vitro incubation with cyanogen bromide or by either in vivo on in vitro incubation with a bacterial methionine N-terminal peptidase (EPO Publ. No. 219 237).

Fusion proteins provide an alternative to direct expression. Usually, a DNA sequence encoding the N-terminal portion of an endogenous bacterial protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the bacteriophage lambda cell gene can be linked at the 5' terminus of a foreign gene and expressed in bacteria. The resulting fusion protein preferably retains a site for a processing enzyme (factor Xa) to cleave the bacteriophage protein from the foreign gene, Nagai et al. (1984) Nature 309:810. Fusion proteins can also be made with sequences from the lacz, Jia et al. (1987) Gene 60:197, trpE, Allen et al. (1987) J. Biotechnol. 5:93; Makoff et al. (1989) J. Gen. Microbiol. 135:11, and EPO Publ. No. 324 647, genes. DNA sequence at the junction of the two amino acid sequences may or may not encode a cleavable site. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that preferably retains a site for a processing enzyme (e.g. ubiquitin specific processingprotease) to cleave the ubiquitin from the foreign protein. Through this method, native foreign protein can be isolated. Miller et al. (1989) Bio/Technology 7:698.

Alternatively, foreign proteins can also be secreted from the cell by creating chimeric DNA molecules that encode a fusion protein comprised of a signal peptide sequence fragment that provides for secretion of the foreign protein in bacteria, U.S. 4,336,336. The signal sequence fragment usually encodes a signal peptide comprised of

WO 93/18150 PCT/EP93/00472

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hydrophobic amino acids which direct the secretion of the protein from the cell. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). Preferably there are processing sites, which can be cleaved either <u>in vivo</u> or <u>in vitro</u> encoded between the signal peptide fragment and the foreign gene.

DNA encoding suitable signal sequences can be derived from genes for secreted bacterial proteins, such as the <u>E. coli</u> outer membrane protein gene (ompA). Masui et al. (1983), in: Experimental Manipulation of Gene Expression; Ghrayeb et al. (1984) EMBO J. 3:2437 and the <u>E. coli</u> alkaline phosphatase signal sequence (phoA), Oka et al. (1985) Proc. Natl. Acad. Sci. 82:7212. As an additional example, the signal sequence of the alpha-amylase gene from various <u>Bacillus</u> strains can be used to secrete heterologous proteins from <u>B. subtilis</u>. Palva et al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EPO Publ. No. 244 042.

Usually, transcription termination sequences recognized by bacteria are regulatory regions located 3' to the translation stop codon, and thus together with the promoter flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Transcription termination sequences frequently include DNA sequences of about 50 nucleotides capable of forming stem loop structures that aid in terminating transcription. Examples include transcription termination sequences derived from genes with strong promoters, such as the trp gene in E. coli as well as other biosynthetic genes.

Usually, the above-described components, comprising a promoter, signal sequence (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (e.g., plasmids) capable of stable maintenance in a host, such as bacteria. The replicon will have a replication system, thus allowing it to

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be maintained in a procaryotic host either for expression or for cloning and amplification. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably contain at least about 10, and more preferably at least about 20 plasmids. Either a high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host.

Alternatively, the expression constructs can be integrated into the bacterial genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to the bacterial chromosome that allows the vector to integrate. Integrations appear to result from recombinations between homologous DNA in the vector and the bacterial chromosome. For example, integrating vectors constructed with DNA from various Bacillus strains integrate into the Bacillus chromosome (EPO Publ. No. 127 328). Integrating vectors may also be comprised of bacteriophage or transposon sequences.

Usually, extrachromosomal and integrating expression constructs may contain selectable markers to allow for the selection of bacterial strains that have been transformed. Selectable markers can be expressed in the bacterial host and may include genes which render bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin (neomycin), and tetracycline. Davies et al. (1978) Annu. Rev.Microbiol. 32:469. Selectable markers may also include biosynthetic genes, such as those in the histidine, tryptophan, and leucine biosynthetic pathways.

Alternatively, some of the above-described components can be put together in transformation vectors. Transformation vectors are usually comprised of a selectable marker that is either maintained in a replicon or developed into an integrating vector.

Expression and transformation vectors, either extra-chromosomal replicons or integrating vectors, have

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PCT/EP93/00472

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been developed for transformation into many bacteria. For example, expression vectors have been developed for, interalia, the following bacteria: Bacillus subtilis, Palv et al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EPO Publ. Nos. 036 259 and 063 953; PCT Publ. No. WO 84/04541; E. coli, Shimatake et al. (1981) Nature 292:128; Amann et al. (1985) Gene 40:183; Studier et al. (1986) J. Mol. Biol. 189:113; EPO Publ. Nos. 036 776, 136 829 and 136 907; Streptococcus cremoris, Powell et al. (1988) Appl. Environ. Microbiol. 54:655; Streptococcus lividans, Powell et al. (1988) Appl. Environ. Microbiol. 54:655; and Streptomyces lividans, U.S. 4,745,056.

Methods of introducing exogenous DNA into bacterial hosts are well-known in the art, and usually include either the transformation of bacteria treated with 15 CaCl, or other agents, such as divalent cations and DMSO. DNA can also be introduced into bacterial cells by electroporation. Transformation procedures usually vary with the bacterial species to be transformed. See, e.g., Masson et al. (1989) FEMS Microbiol. Lett. 60:273; Palva et 20 al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EPO Publ. Nos. 036 259 and 063 953; PCT Publ. No. WO 84/04541, for Miller et al. (1988) Proc. Natl. Acad. Sci. Bacillus; 85:856; Wang et al. (1990) J. Bacteriol. 172:949, for Campylobacter; Cohen et al. (1973) Proc. Natl. Acad. Sci. 25 69:2110; Dower et al. (1988) Nucleic Acids Res. 16:6127; Kushner (1978) "An improved method for transformation of E. coli with ColE1-derived plasmids," In Genetic Engineering: Proceedings of the International Symposium on Genetic Engineering (eds. H.W. Boyer and S. Nicosia); Mandel et al. 30 (1970) J. Mol. Biol. 53:159; Taketo (1988) Biochim. Biophys. Acta 949:318, for Escherichia; Chassy et al. (1987) FEMS Microbiol. Lett. 44:173, for Lactobacillus; Fiedler et al. (1988) Anal. Biochem 170:38, for <u>Pseudomonas</u>; Augustin et al. (1990) FEMS Microbiol. Lett. 66:203, for Staphylococcus; 35 Barany et al. (1980) J. Bacteriol. 144:698; Harlander (1987) "Transformation of Streptococcus lactis by electroporation, in: Streptococcal Genetics (ed. J. Ferretti and R. Curtiss III); Perry et al. (1981) Infec. Immun. 32:1295; Powell et

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al. (1988) Appl. Environ. Microbiol. 54:655; Somkuti et al. (1987) Proc. 4th Evr. Cong. Biotechnology 1:412, for Streptococcus.

iv. Yeast Expression

Yeast expression systems are also known to one of ordinary skill in the art. A yeast promoter is any DNA sequence capable of binding yeast RNA polymerase and initiating the downstream (3') transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site (the "TATA Box") and a transcription initiation site. A yeast promoter may also have a second domain called an upstream activator sequence (UAS), which, if present, is usually distal to the structural gene. The UAS permits regulated (inducible) expression. Constitutive expression occurs in the absence of a UAS. Regulated expression may be either positive or negative, thereby either enhancing or reducing transcription.

Yeast is a fermenting organism with an active metabolic pathway, therefore sequences encoding enzymes in the metabolic pathway provide particularly useful promoter Examples include alcohol dehydrogenase (ADH) sequences. (EPO Publ. No. 284 044), enolase, glucokinase, glucose-6phosphate isomerase, glyceraldehyde-3-phosphatedehydrogenase (GAP GAPDH), or hexokinase, phosphofructokinase, 3-phosphoglycerate mutase, and pyruvate kinase (PyK) (EPO Publ. No. 329 203). The yeast PHO5 gene, encoding acid phosphatase, also provides useful promoter sequences, Myanohara et al. (1983) Proc. Natl. Acad. Sci. USA 80:1.

In addition, synthetic promoters which do not occur in nature also function as yeast promoters. For example, UAS sequences of one yeast promoter may be joined with the transcription activation region of another yeast promoter, creating a synthetic hybrid promoter. Examples of such hybrid promoters include the ADH regulatory sequence linked to the GAP transcription activation region (U.S.

WO 93/18150 PCT/EP93/00472

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4,876,197 and U.S. 4,880,734). Other examples of hybrid promoters include promoters which consist of the regulatory sequences of either the ADH2, GAL4, GAL10, or PHO5 genes, combined with the transcriptional activation region of a glycolytic enzyme gene such as GAP or PyK (EPO Publ. No. 164 556). Furthermore, a yeast promoter can include naturally occurring promoters of non-yeast origin that have the ability to bind yeast RNA polymerase and initiate transcription. Examples of such promoters include, inter alia, Cohen et al. (1980) Proc. Natl. Acad. Sci. USA 77:1078; Henikoff et al. (1981) Nature 283:835; Hollenberg et al. (1981) Curr. Topics Microbiol. Immunol. 96:119; Hollenberg et al. (1979) "The Expression of Bacterial Antibiotic Resistance Genes in the Yeast Saccharomyces cerevisiae," in: Plasmids of Medical, Environmental and Commercial Importance (eds. K.N. Timmis and A. Puhler); Mercerau-Puigalon et al. (1980) Gene 11:163; Panthier et al. (1980) Curr. Genet. 2:109.

A DNA molecule may be expressed intracellularly in yeast. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by in vitro incubation with cyanogen bromide.

Fusion proteins provide an alternative for yeast expression systems, as well as in mammalian, baculovirus, and bacterial expression systems. Usually, a DNA sequence encoding the N-terminal portion of an endogenous yeast protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the yeast or human superoxide dismutase (SOD) gene, can be linked at the 5' terminus of a foreign gene and expressed in yeast. The DNA sequence at the junction of the two amino acid sequences may or may not encode a cleavable site. See e.g., EPO Publ. No. 196 056. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that

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preferably retains a site for a processing enzyme (e.g. ubiquitin-specific processing protease) to cleave the ubiquitin from the foreign protein. Through this method, therefore, native foreign protein can be isolated (see, e.g., PCT Publ. No. WO 88/024066).

Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provide for secretion in yeast of the foreign protein. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either in vivo or in vitro. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell.

DNA encoding suitable signal sequences can be derived from genes for secreted yeast proteins, such as the yeast invertase gene (EPO Publ. No. 012 873; JPO Publ. No. 62,096,086) and the A-factor gene (U.S. 4,588,684). Alternatively, leaders of non-yeast origin, such as an interferon leader, exist that also provide for secretion in yeast (EPO Publ. No. 060 057).

A preferred class of secretion leaders are those that employ a fragment of the yeast alpha-factor gene, which contains both a "pre" signal sequence, and a "pro" region. The types of alpha-factor fragments that can be employed include the full-length pre-pro alpha factor leader (about 83 amino acid residues) as well as truncated alpha-factor leaders (usually about 25 to about 50 amino acid residues) (U.S. 4,546,083 and U.S. 4,870,008; EPO Publ. No. 324 274). Additional leaders employing an alpha-factor leader fragment that provides for secretion include hybrid alpha-factor leaders made with a presequence of a first yeast, but a proregion from a second yeast alphafactor. (See e.g., PCT Publ. No. WO 89/02463.)

Usually, transcription termination sequences recognized by yeast are regulatory regions located 3' to the translation stop codon, and thus together with the promoter

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PCT/EP93/00472

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flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminator sequence and other yeast-recognized termination sequences, such as those coding for glycolytic enzymes.

Usually, above-described the components, comprising a promoter, leader (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs maintained often in a replicon, such as an extrachromosomal element (e.g., plasmids) capable of stable maintenance in a host, such as yeast or bacteria. replicon may have two replication systems, thus allowing it to be maintained, for example, in yeast for expression and in a procaryotic host for cloning and amplification. Examples of such yeast-bacteria shuttle vectors include YEp24, Botstein et al. (1979) Gene 8:17-24; pCl/1, Brake et al. (1984) Proc. Natl. Acad. Sci USA 81:4642-4646; and YRp17, Stinchcomb et al. (1982) J. Mol. Biol. 158:157. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably have at least about 10, and more preferably at least about 20. A high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host.

Alternatively, the expression constructs can be integrated into the yeast genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to a yeast chromosome that allows the vector to integrate, and preferably contain two homologous sequences flanking the expression construct. Integrations appear to result from recombinations between homologous DNA in the vector and the yeast chromosome, Orr-Weaver et al. (1983) Methods in Enzymol. 101:228-245. An integrating vector may be directed to a specific locus in yeast by selecting the appropriate homologous sequence for inclusion in the vector. One or more expression construct may integrate, possibly

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affecting levels of recombinant protein produced, Rine et al. (1983) Proc. Natl. Acad. Sci. USA 80:6750. The chromosomal sequences included in the vector can occur either as a single segment in the vector, which results in the integration of the entire vector, or two segments homologous to adjacent segments in the chromosome and flanking the expression construct in the vector, which can result in the stable integration of only the expression construct.

10 Usually, extrachromosomal integrating and expression constructs may contain selectable markers to allow for the selection of yeast strains that have been transformed. Selectable markers may include biosynthetic genes that can be expressed in the yeast host, such as ADE2, 15 HIS4, LEU2, TRP1, and ALG7, and the G418 resistance gene, which confer resistance in yeast cells to tunicamycin and G418, respectively. In addition, a suitable selectable marker may also provide yeast with the ability to grow in the presence of toxic compounds, such as metal. For example, the presence of CUP1 allows yeast to grow in the 20 presence of copper ions. Butt et al. (1987) Microbiol, Rev. 51:351.

Alternatively, some of the above-described components can be put together into transformation vectors. Transformation vectors are usually comprised of a selectable marker that is either maintained in a replicon or developed into an integrating vector.

Expression and transformation vectors, either extrachromosomal replicons or integrating vectors, have been developed for transformation into many yeasts. For example, expression vectors have been developed for, inter alia, the following yeasts: Candida albicans, Kurtz, et al. (1986) Mol. Cell. Biol. 6:142; Candida maltosa, Kunze, et al. (1985) J. Basic Microbiol. 25:141; Hansenula polymorpha, Gleeson, et al. (1986) J. Gen. Microbiol. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302; Kluyveromyces fraqilis, Das, et al. (1984) J. Bacteriol. 158:1165; Kluyveromyces lactis, De Louvencourt et al. (1983) J. Bacteriol. 154:737; Van den Berg et al. (1990)

Bio/Technology 8:135; Pichia quillerimondii, Kunze et al. (1985) J. Basic Microbiol. 25:141; Pichia pastoris, Cregg, et al. (1985) Mol. Cell. Biol. 5:3376; U.S. 4,837,148 and U.S. 4,929,555; Saccharomyces cerevisiae, Hinnen et al. (1978) Proc. Natl. Acad. Sci. USA 75:1929; Ito et al. (1983) J. Bacteriol. 153:163; Schizosaccharomyces pombe, Beach et al. (1981) Nature 300:706; and Yarrowia lipolytica, Davidow, et al. (1985) Curr. Genet. 10:380471 Gaillardin, et al. (1985) Curr. Genet. 10:49.

10 Methods of introducing exogenous DNA into yeast hosts are well-known in the art, and usually include either the transformation of spheroplasts or of intact yeast cells treated with alkali cations. Transformation procedures usually vary with the yeast species to be transformed. See 15 e.g., Kurtz et al. (1986) Mol. Cell. Biol. 6:142; Kunze et al. (1985) J. Basic Microbiol. 25:141, for Candida; Gleeson et al. (1986) J. Gen. Microbioy. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302, for Hansenula; Das et al. (1984) J. Bacteriol. 158:1165; De Louvencourt et al. (1983) 20 J. Bacteriol. 154:1165; Van den Berg et al. Bio/Technology 8:135, for <u>Kluyveromyces</u>; Cregg et al. (1985) Mol. Cell. Biol. 5:3376; Kunze et al. (1985) J. Basic Microbiol. 25:141; U.S. 4,837,148 and U.S. 4,929,555, for Pichia; Hinnen et al. (1978) Proc. Natl. Acad. Sci. USA 75;1929; Ito et al. (1983) J. Bacteriol. 153:163, for 25 Saccharomyces; Beach et al. (1981) Nature 300:706, for Schizosaccharomyces; Davidow et al. (1985) Curr. Genet. 10:39; Gaillardin et al. (1985) Curr. Genet. 10:49, for Yarrowia.

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E. Vaccines

Each of the <u>H. pylori</u> proteins discussed herein may be used as a sole vaccine candidate or in combination with one or more other antigens, the latter either from <u>H. pylori</u> or other pathogenic sources. Preferred are "cocktail" vaccines comprising, for example, the cytotoxin (CT) antigen, the CAI protein, and the urease. Additionally, the hsp can be added to one or more of these components. These vaccines may either be prophylactic (to

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prevent infection) or therapeutic (to treat disease after infection).

Such vaccines comprise H. pylori antigen or antigens, usually in combination with "pharmaceutically acceptable carriers", which include any carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition. Suitable carriers are typically large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, lipid aggregates (such as oil droplets or liposomes), and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Additionally, these carriers may function immunostimulating as agents ("adjuvants"). Furthermore, the antigen may be conjugated to a bacterial toxoid, such as a toxoid from diphtheria, tetanus, cholera, H. pylori, etc. pathogens.

Preferred adjuvants to enhance effectiveness of the composition include, but are not limited to: aluminum salts (alum), such as aluminum hydroxide, aluminum phosphate, aluminum sulfate, etc; (2) oil-in-water emulsion formulations (with without _ other or immunostimulating agents such as muramyl peptides (see below) or bacterial cell wall components), such as for example (a) MF59 (PCT Publ. No. WO 90/14837), containing 5% Squalene, 0.5% Tween 80, and 0.5% Span 85 (optionally containing various amounts of MTP-PE (see below), although not required) formulated into submicron particles using a microfluidizer such as Model 110Y microfluidizer (Microfluidics, Newton, MA), (b) SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-blocked polymer L121, and thr-MDP (see below) either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion, and (c) RibiTM adjuvant system (RAS), (Ribi Immunochem, Hamilton, MT) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably (DetoxTM); (3) saponin adjuvants, such as MPL + CWS

StimulonTM (Cambridge Bioscience, Worcester, MA) may be used or particles generated therefrom such as ISCOMs (immunostimulating complexes); (4) Complete Freunds Adjuvant (CFA) and Incomplete Freunds Adjuvant (IFA); (5) cytokines, such as interleukins (IL-1, IL-2, etc.), macrophage colony stimulating factor (M-CSF), tumor necrosis factor (TNF), etc; and (6) other substances that act as immunostimulating agents to enhance the effectiveness of the composition. Alum and MF59 are preferred.

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As mentioned above, muramyl peptides include, but are not limited to, N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-L-alanyl-D-isoglutamine (nor-MDP), N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-huydroxyphosphoryloxy)-ethylamine (MTP-PE), etc.

The immunogenic compositions (e.g., the antigen, pharmaceutically acceptable carrier, and adjuvant) typically will contain diluents, such as water, saline, glycerol, ethanol, etc. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles.

Typically, the immunogenic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection may also be prepared. The preparation also may be emulsified or encapsulated in liposomes for enhanced adjuvant effect, as discussed above under pharmaceutically acceptable carriers.

Immunogenic compositions used as vaccines comprise an immunologically effective amount of the antigenic polypeptides, as well as any other of the above-mentioned components, as needed. By "immunologically effective amount", it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, the taxonomic group of individual to be treated (e.g., nonhuman primate,

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primate, etc.), the capacity of the individual's immune system to synthesize antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

The immunogenic compositions are conventionally administered parenterally, e.g., by injection, either subcutaneously or intramuscularly. Additional formulations suitable for other modes of administration include oral and pulmonary formulations, suppositories, and transdermal applications. Oral formulations are most preferred for the H. pylori proteins. Dosage treatment may be a single dose schedule or a multiple dose schedule. The vaccine may be administered in conjunction with other immunoregulatory agents.

F. Immunodiagnostic Assays

H. pylori antigens can be used in immunoassays to detect antibody levels (or conversely H. pylori antibodies can be used to detect antigen levels) and correlation can be made with gastroduodenal disease and with duodenal ulcer in particular. Immunoassays based on well defined, recombinant antigens be can developed to replace invasive the diagnostics methods that are used today. Antibodies to H. pylori proteins within biological samples, including for example, blood or serum samples, can be detected. Design of the immunoassays is subject to a great deal of variation, and a variety of these are known in the art. Protocols for the immunoassay may be based, for example, upon competition, or direct reaction, or sandwich type assays. Protocols may also, for example, use solid supports, or may be by immunoprecipitation. Most assays involve the use of labeled antibody or polypeptide; the labels may be, for example, fluorescent, chemiluminescent, radioactive, molecules. Assays which amplify the signals from the probe are also known; examples of which are assays which utilize biotin and avidin, and enzyme-labeled and mediated immunoassays, such as ELISA assays.

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Kits suitable for immunodiagnosis and containing the appropriate labeled reagents are constructed by packaging the appropriate materials, including the compositions of the invention, in suitable containers, along with the remaining reagents and materials (for example, suitable buffers, salt solutions, etc.) required for the conduct of the assay, as well as suitable set of assay instructions.

G. Examples

The examples presented below are provided as a further guide to the practitioner of ordinary skill in the art and are not to be construed as limiting the invention in any way.

i. H. pylori cytotoxin (CT) antigen

15 <u>1. Materials and methods</u>

For general materials and methods relating to \underline{H} . \underline{Pylori} growth and DNA isolation, see sections ii and iii below, relating to CAI antigen and hsp, respectively.

a. Cloning

Two mixtures of degenerate oligonucleotides were synthesized using an Applied Biosystems model 380B DNA synthesizer. These mixtures were used at a concentration of 4 micromolar in a 100 microliter polymerase chain reaction with 200 nanograms of purified DNA using the Genamp PCR kit according to the manufacturers instructions. The reaction was incubated for 1 minute at 94 degrees centigrade, 2 minutes at 48 degrees centigrade and 2 minutes at 56 degrees centigrade. The reaction mix was subjected to 30 cycles of these conditions.

Analysis of the products of this reaction by agarose gel electrophoresis revealed a prominent approximately 87 bp DNA fragment. After digestion with the restriction enzymes XbaI and EcoRI, the fragment was ligated to the Bluscript SK+ (Stratgene) plasmid which had previously also been digested with XbaI and EcoRI. The ligation mixture was used to transform competent E. coli by electroporation at 2000V and 25 microfarads using (200 Ω) a BioRad Gene Pulser (California). Transformed E. coli were selected for growth on L-agar plates containing 100

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micrograms per milliliter ampicillin. Plasmid DNA was extracted from positive <u>E.coli</u> isolates and subjected to sequence analysis using the Sequenase 2 (United States Biochemical Corporation) DNA sequencing kit according to the manufacturers instructions.

b. Preparation of libraries

(1) Library of HindIII fragments

Seven micrograms of purified DNA were digested to completion with the restriction enzyme HindII. micrograms of Bluescript SK+ plasmid DNA were digested to completion with HindIII then treated with calf intestinal phosphatase. Both DNA mixtures were purified by agitation with a water saturated phenol then precipitated by addition of ethyl alcohol to 67% V/V. Both DNAs were resuspended in 50 microliters of water. 0.7 micrograms of DNA fragments were mixed with 0.3 micrograms of Bluescript DNA in 50 microliters of a solution containing 25 mM Tris ph 7.5, 10mM MgCl2 and 5 units of T4 DNA ligase. This mix was incubated at 15 deg. centigrade for 20 hours after which the DNA was extracted with water saturated phenol and precipitated from ethyl alcohol. The DNA was subsequently resuspended in 50 microL. of water. Introduction of 1 microL of this DNA into E.coli by eletroporation resulted in approximately 3000-10,000 ampicillin resistant bacterial colonies.

2) Library of EcoRI fragments.

About 0.7 microg. of EcoRI digested DNA was purified and mixed with 0.45 micrograms of Bluescript SK+ plasmid which had been previously digested with EcoRI and treated with calf intestinal phosphatase. The fragments were ligated in 50 microL of solution. After purification and precipitation, the DNA was resuspended in 50 microL of water. Electroporation of E. coli with 1 microL of this solution resulted in approximately 200 ampicillin resistant bacterial colonies.

In order to identify suitable restriction fragments from the genome for further cloning, the plasmid was uniformly labeled with 32p and used as a probe to analyze DNA from the strain CCUG digested with various restriction enzymes, separated on agarose gel

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electrophoresis and transferred to nitrocellulose filter. The probe revealed a unique approximately 3.5kb HindIII restriction fragment. A library of HindIII digested DNA fragments was prepared and cloned in the Bluescript plasmid vector. This library was screened with 32p labeled DNA corresponding to the 87 bp fragment previously cloned. Two clones containing identical approximately 3.3 kbp hindIII fragments were identified. DNA sequencing of these HindIII fragments revealed sequences capable of coding for the 23 amino acids corresponding to the amino terminus of the previously described 87 kDa cytotoxin. These sequences comprised part of an open reading frame of approximately 300 nucleotides which terminated at the extremity of the fragment delimited by a HindIII restriction site. The sequence also revealed the existence of an EcoRI restriction site within the putative open reading frame 120 bp away from the HindIII site.

A 32p labeled probe corresponding to the sequences between the EcoRI site and the HindIII site was used to screen a library of EcoR fragments from DNA cloned in the Bluescript SK vector. This probe revealed two clones containing approximately 7.3 kbp fragments. DNA sequencing of these fragments revealed a continuous open reading frame which overlapped with the sequences determined from the 3.2 kbp HindIII fragments. The DNA sequence of these overlapping fragments and the conceptual translation of the single long open reading frame contained are shown in Figs. 1 and 2, respectively.

It should be noted that these clones were found to be extremely unstable. The initial colonies identified in the screening were so small as to be difficult to detect. Expansion of these clones by traditional methods of subculturing for 16-18 hours resulted in very heterogeneous populations of plasmids due to DNA rearrangement and deletion. Sufficient quantities of these clones were grown by subculturing for 8-10 hours in the absence of antibiotic selection. In this fashion, although yields of plasmid were relatively low, selection and outgrowth of bacteria containing viable rearranged plasmid were avoided.

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c. Screening of DNA libraries

The product of the PCR reaction which contained the predominant 87 bp fragment was labeled with 32p by the random priming method using the Prime-a-gene kit (Promega). This labeled probe was used in a hybridization reaction with DNA from approximately 3000 bacterial clones immobilized on nitrocellulose filters. The hybridization reaction was carried out at 60 degrees centigrade in a solution of 0.3M NaCl. A positive bacterial clone was expanded and plasmid DNA was prepared. The plasmid contained an insert of approximately 3.3kb of DNA and was designated TOXHH1.

A 120 bp fragment containing the sequences between position 292 and 410 shown in Fig. 1 was derived from the plasmid TOXHH1 and used to screen approximately 400 colonies of the library of EcoRI fragments. A positive clone was isolated which contained approximately 7.3kb of DNA sequences and was designated TOXEE1.

The nucleotide sequence shown in Fig. 1 was derived from the clones TOXHH1 and TOXEE1 using the Sequenase 2 sequencing kit. The nucleotides between position 1 and 410 in Fig. 1 were derived from TOXHH1 and those between 291 and 3507 were derived from TOXEE1. E. coli containing plasmids TOXHH1 and TOXEE1 have been deposited with the American Type Culture Collection, see below.

d. Preparation of antisera against the cytotoxin

A DNA fragment corresponding to nucleotides 116-413 of the sequence shown in Fig. 1 was cloned into the bacterial expression vector pex 34 A, such that on induction of the bacterial promoter, a fusion protein was produced which contained a part of the MS2 polymerase polypeptide fused to the amino acids of the cytotoxin polypeptide and including the 23 amino acids previously identified. Approximately 200 micrograms of this fusion protein were partially purified by acrylamide gel electrophoresis and used to immunize rabbits by standard procedures.

Antisera from these rabbits taken after 3 immunizations spaced 1 month apart was used to probe protein extracts from a cytotoxin positive and a cytotoxin negative strain of <u>H. pylori</u> in standard immunoblotting experiments.

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The antisera revealed a polypeptide which migrated on denaturing polyacrylamide gel electrophoresis with an apparent molecular mass of 100 kDa. This polypeptide was detected in protein extracts of the cytotoxin positive but not the cytotoxin negative strain. Serum collected prior to immunization did not react with this polypeptide.

e. Partial purification of vacuolating activity

Total <u>H. pylori</u> membranes at a concentration of 6 mg/ml were solubilized in a solution of 1% CHAPS, 0.5 M NaCl, 10 mM Hepes pH 7.4, 2.5 mM EDTA, 20% sucrose for 1 hour at 4°C. This mixture was then applied to a discontinuous sucrose gradient containing steps of 30%, 35%, 40% and 55% sucrose and subjected to ultracentrifugation for 17 hours at 20000 x g. The gradient was fractionated and each fraction was tested for vacuolating activity and for urease activity. Vacuolating activity associated with urease activity was found in several fractions of the gradient. A peak of vacuolating activity was also found in the topmost fractions of the gradient and these fractions were essentially free of urease activity.

This urease-independent vacuolating activity was further fractionated by stepwise precipitation with ammonium sulphate between concentrations of 20% to 34%. Denaturing polyacrylamide gel electrophoresis of the proteins precipitated at different concentrations of ammonium sulphate revealed a predominant polypeptide of about 100 kDa which copurified with the vacuolating activity. This polypeptide was recognised by the rabbit antisera raised against the recombinant fusion protein described above.

30 <u>2. Results</u>

Two overlapping fragments corresponding to about 10 kbp of the <u>H. pylori</u> genome have been cloned. These clones contain a gene consisting of 3960 bp (shown in Fig.1) which is capable of coding for a polypeptide of 1296 amino acids (shown in Fig.2). The molecular weight of this putative polypeptide is 139.8 kd. The nucleotide sequence AGGAAG 9 bp upstream of the methionine codon at position 18 in Fig.1 resembles closely the consensus Shine-Dalgarno sequence and supports the hypothesis that this methionine

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represents the initiator methionine for synthesis of the polypeptide. A 30 bp nucleotide sequence which begins 10 bp downstream of the putative stop codon at position 3906 in Fig. 1 resembles closely the the structure of prokaryotic transcription terminators and is likely to represent the end of the messenger RNA coding sequences.

The cytotoxin gene is defined as coding for a polypeptide precursor of the <u>H. pylori</u> vacuolating activity by the following criteria:

(i) The putative polypeptide contains the 23 amino acid sequence (Fig. 2, positions 34-56) identified as the amino terminus of the previously described 87 kDa vaculating protein, Clover et al., J. Biol. Chem. 267:10570-75 (1992). This sequence is preceded by 33 amino acids which resemble prokaryotic leader sequences; thus, this sequence is likely to represent the amino terminus of a mature protein;

(ii) Rabbit antisera specific for a 100 amino acid fragment of the putative polypeptide containing the proposed amino terminus recognized a 100 kDa polypeptide in a cytotoxin positive but not a cytotoxin negative strain of H. pylori. This 100 kDa polypeptide copurifies with vacuolating activity from H. pylori membranes.

In sum, the gene described herein codes for an approximately 140 kDa polypeptide which is processed to a 100 kDa polypeptide involved in <u>H. pylori</u> cytotoxic activity. The 87 kDa polypeptide previously described must result from either further processing of the 100 kDa polypetide or from proteolytic degradation during purification.

ii. H. pylori CAI antigen

- 1. Materials and methods
- a. Origin of materials

Clones A1, 64/4, G5, A17, 24 and 57/D were obtained from the lambda gt11 library. Clone B1 was obtained from a genomic plasmid library of HindIII fragments. 007 was obtained by PCR. The H. pylori strains producing the cytotoxin were: G10, G27, G29, G32, G33, G39, G56, G65, G105, G113A. The noncytotoxic strains were: G12, G21, G25,

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G47, G50, G204. They were isolated from endoscopy biopsy specimens at the Grosseto Hospital, (Tuscany, Italy). The strain CCUG 17874 (cytotoxin positive), was obtained from the Culture Collection of the University of Gotheborg. The noncytotoxic strains Pylo 2U+ (urease positive) and Pylo 2U-(urease negative) were obtained from F. Megraud, Centre Hospitalier, Bordeaux (France). E. coli strains DH10B (Bethesda Research Laboratories), TG1, K12 delta H1 delta trp, Y1088, Y1089, Y1090 are known in the art. Plasmid Bluescript SK+ (Stratagene, La Jolla, CA) was used as a cloning vector. The pEx34 a, b, c plasmids for the expression of MS2 fusion proteins have been previously described. The lambda gt11 phage vector used for the expression library is from the lambda gt11 cloning system kit (Bethesda Research Laboratories). E. coli strains were cultured in LB medium (24). H. pylori strains were plated onto selective media (5% horse blood, Columbia agar base with Dent or Skirrow's antibiotic supplement, 0.2% cyclodextrin) or in Brucella broth liquid medium containing 5% fetal bovine serum (6) or 0.2% cyclodextrin (25).

b. Growth of <u>H. pylori</u> and DNA isolation

H. pylori strains were cultured in solid or liquid media for 3 days at 37 °C, both in microaerophilic atmosphere using Oxoid (Basingstoke, England) or Becton and Dickinson (Cockeysville, MD) gas pack generators or in an incubator containing air supplemented with 5% CO2, (26). The bacteria were harvested and resuspended in STE (NaCl 0.1M, Tris-HCl 10mM pH 8, EDTA 1 mM pH 8) containing lysozyme at a final concentration of 100 micrograms/ml and incubated at room temperature for 5 min. To lyse the bacteria SDS was added to a final concentration 1% and heated at 65 °C. After the addition of proteinase K at final concentration of 25 micrograms/ml the solution was incubated at 50° for 2 hours. The DNA was purified by CsCl gradient in the presence of ethidium bromide, precipitated with 77% ethanol and recovered with a sealed glass capillary.

c. Construction and screening of a lambda gt11 expression library

To generate the lambda gtll expression library,

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genomic DNA from the CCUG 17874 strain partially digested with the restriction enzymes HaeIII and AluI was used. After fractionation on 0.8% agarose gel, the DNA between 0.6 and 8 Kb in size was eluted using a Costar Spin-X (0.22 micron) microcentrifuge filter. The products from each digestion were combined, and used to construct the expression library, using the lambda gtll cloning system kit (Bethesda Research Laboratories) and the Gigapack II Gold packaging kit (Stratagene, La Jolla, CA). The library that contained 0.8-1 x 106 recombinant phages was amplified in E. coli Y1088, obtaining 150 ml of a lysate with a titer of 109 phages/ml, 85% of which were recombinant and had an average insert size of 900 base pairs,. Immunological screening was performed by standard procedures, using the Protoblot system (Promega, Madison, WI).

d. Construction of plasmid libraries

Attempts to make complete genomic libraries of partially digested chromosomal DNA, using standard vectors such as EMBL4 or lambda Dash encountered the difficulties described also by many authors in cloning <u>H. pylori</u> DNA and failed to give satisfactory libraries. Therefore, partial libraries were obtained using genomic DNA from strains CCUG 17874, G39 and G50 digested with the restriction enzyme HindIII, cloned in the Bluescript SK+. DNA ligation, electroporation of <u>E. coli</u> DH 10B, screening, and library amplification have been performed. Libraries ranging from 70000 to 85000 colonies with a background not exceeding the 10% were obtained.

2. DNA manipulation and nucleotide sequencing

DNA manipulation was performed using standard procedures. DNA sequencing was performed using Sequenase 2.0 (USB) and the DNA fragments shown in Fig. 3 subcloned in Bluscript KS+. Each strand was sequenced at least three times. The region between nucleotides 1533 and 2289, for which a DNA clone was not available, was amplified by PCR and sequenced using asymmetric PCR, and direct sequencing of amplified products. The overlapping of this region, was confirmed by one and double side anchored PCR: an external universal anchor (5'-GCAAGCTTATCGATGTCGAGCT-3'/ 5'-

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GACTCGAGTCGACATCGA-3') containing a protruding 5' HindIII sequence, and the recognition sites of ClaI, SalI, XhoI, was ligated to primer-extended DNA and amplified. A second round of PCR using nested primers was then used to obtain fragments of DNA suitable for cloning and sequencing. DNA sequence data were assembled and analyzed with the GCG package (Genetics Computer Group, Inc., Madison, WI) running on a VAX 3900 under VMS. The GenBank and EMBL databases were examined using the EMBL VAXcluster.

f. Protein preparation and ELISA 10

Protein extracts were obtained by treating H. pylori pellets with 6 M guanidine. Western blotting, SDS-PAGE, electroelution were performed by standard procedures. Fusion proteins were induced and purified by electrocution or by ion exchange chromatography. Purified proteins were used to immunize rabbits and to coat microtiter plates for ELISA assays. Sera from people with normal mucosa, blood donors and patients were obtained from A. Ponzetto (Torino, Italy) Clinical diagnosis was based on histology of gastric biopsies. Vacuolating activity of samples was tested on HeLa cells as described by Cover et al. Infect. Immun. 59:1264-70 (1991).

2. Results

Immunodominance and cytotoxicity a.

25 Western blots of <u>H. pylori</u> guanidine extracts probed with sera from patients with gastroduodenal disease showed that a protein of 130 kDa that is a minor component in the Coomassie blue stained gel was strongly recognized by all sera tested. The CAI protein was electroeluted and used 30 to raise a mouse serum that in a Western blot recognized only this protein. This serum was then used to detect by Western blotting the CAI protein in extracts of the H. <u>pylori</u> strains. The antigen was present in the all 10 strains that had vacuolizing activity on HeLa cells while it was absent in the eight strains that did not have such activity; in addition, the size of the protein varied slightly among the strains. The CAI antigen was not detected by western blotting in the other species tested such as Campylobacter jejuni, Helicobacter mustelae, E. coli, and

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Bordetella pertussis.

b. Structure of the cai gene

were screened using the mouse serum specific for the CAI antigen and with a pool of sera from patients with gastroduodenal diseases. The mouse serum detected positive clones at a frequency of 3 x 10⁻³. Sequence analysis of 8 clones revealed that they were all partially overlapping with clone A1 shown in Fig. 3. The pool of human sera identified many clones containing different regions of the cai gene, including clones 57/D, 64/4 and 24 and several clones overlapping clone A1.

In Fig. 3, clones A1, 64/4, G5, A17, 24, and 57/D were obtained from the lambda gt11 library. Clone B1 was obtained from a plasmid library of HindIII fragments. E. coli containing plasmids 57/D, 64/4, B1 (B/1), and P1-24 (the latter most plasmid from nucleotide 2150 to 2650) have been deposited with the American Type Culture Collection (ATCC), see below. 007 was obtained by PCR. reading frame is shown at the bottom of Fig. 3. indicate the position and direction of the synthetic oligonucleotides used as primers for sequencing, and the position of insertion of the repeated sequence of G39 is shown. The nucleotide and amino acid sequence of one of the repeated sequences found in strain G39 is also shown. The capital letters indicate the sequences D1, D2, and D3 duplicated from the cai gene, the small letters indicate the nucleotide and amino acid linkers, P=promoter, T=terminator.

The nucleotide sequence of the entire region was determined using the clones derived from the lambda gt11 library, the clone B1 isolated from the HindIII plasmid library, and the fragment 007 that was obtained by PCR of the chromosomal DNA. Computer analysis of the 5925 nucleotide sequence revealed a long open reading frame spanning nucleotides 535 to 3977 that was in frame with the fusion proteins deriving from the lambda gt11 clones 64/4, 24 and A1 and A17. Clone 57/D contained an open reading frame only in the 3' end of cloned fragment and therefore

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could not make a gene fusion with the beta galactosidase gene of lambda gt11. The presence of an immunoreactive protein in the lambda gt11 clone 57/D could only be explained by the presence of an endogenous promoter driving the expression of a non fused protein. This hypothesis was proven to be true by subcloning in both direction the insert 57/4 into the Bluescript plasmid vector and showing that an immunoreactive protein was obtained in both cases. A conclusive evidence that the gene identified was indeed coding for the CAI antigen was obtained by subcloning the inserts A17 and 64/4 in the pEx 34B plasmid vectors to obtain fusion proteins that were purified and used to immunize rabbits. The sera obtained, recognized specifically the CAI antigen band in cytotoxic H. pylori strains.

The cai gene coded for a putative protein of 1147 amino acids, with predicted molecular weight of 128012.73 Daltons and an isoelectric point of 9.72. The basic properties of the purified protein were confirmed by two dimensional gel electrophoresis. The codon usage and the GC content (37%) of the gene were similar to that described for other <u>H. pylori</u> genes (13,26). A putative ribosome binding site: AGGAG, was identified 5 base pairs_upstream from the proposed ATG starting codon. Computer search for promoter sequences of the region upstream from the ATG start codon, identified sequences resembling either -10 or - 35 regions, however, a region with good consensus to an E. coli promoter, or resembling published H. pylori promoter sequences was not found. Primer extension analysis of purified <u>H. pylori</u> RNA showed that 104 and 214 base pairs upstream from the ATG start codon there are two transcriptional start sites. Canonical promoters could not upstream from either transcriptional be identified initiation sites. The expression of a portion of the CAI antigen by clone 57/D suggests that E. coli is also recognizing a promoter in this region, however, it is not clear whether E. coli recognizes the same promoters of H. pylori or whether the <u>H. pylori</u> DNA that is rich in A-T provides E. coli with regions that may act as promoters. A rho independent terminator was identified downstream from

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the stop codon. In Fig. 4, the AGGAG ribosome binding site and terminator are underlined, and the repeated sequence and motif containing 6 asparagines are boxed. The CAI antigen was very hydrophilic, and did not show obvious leader peptide or transmembrane sequences. The most hydrophilic region was from amino acids 600 to 900, where also a number of unusual features can be observed: the repetition of the sequences EFKNGKNKDFSK and EPYIA, and the presence of a stretch of six contiguous asparagines (boxed in Fig. 4).

10 c. Diversity of the cai gene

Diversity of the gene appears to be generated by internal duplications. To find out the mechanism of size heterogeneity of the CAI proteins in different strains, the structure of one of the strains with a larger CAI protein (G39) was analyzed using Southern blotting, PCR and DNA sequencing. The results showed that the cai gene of G39 and CCUG 17874 were identical in size until position 3406, where the G39 strain was found to contain an insertion of 204 base pairs, made by two identical repeats of 102 base pairs. Each repeat was found to contain sequences deriving from the duplication of 3 segments of DNA (sequences D1, D2 and D3 in Fig. 3) coming from the same region of the cai gene and connected by small linker sequences. A ·schematic representation of the region where the insertion occurred and of the insertion itself is shown in Fig. 3.

d. cai gene absent in noncytotoxic strains

To investigate why the CAI antigen was absent in the noncytotoxic strains, DNA from two of them (G50 and G21), was digested with EcoRI, HindIII and HaeIII restriction enzymes, and tested by Southern blotting using two probes internal to the <u>cai</u> gene, spanning nucleotides 520-1840 and 2850-4331 respectively. Both probes recognized strongly hybridizing bands in strains CCUG 17874 and G39. The bands varied in size in the two strains, in agreement with the gene diversity. However, neither probe hybridized the G50 and G21 DNA. This showed that the noncytotoxic strains tested do not contain the <u>cai</u> gene.

e. Serum antibodies

The presence of serum antibodies against the CAI

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antigen correlated with gastroduodenal diseases. To study the quantitative antibody response to the CAI antigen, the fusion protein produced by the A17 fragment subcloned in pEx34 was purified to homogeneity and used to coat microtiter plates for an ELISA test. In this assay, the patients with gastroduodenal pathologies had an average ELISA titer that was significantly higher than that found in randomly selected blood donors and people with normal gastric mucosa. To evaluate whether the antibody titer correlated with a particular gastroduodenal disease, the sera from patients with known histological diagnosis were tested in the ELISA assay. Patients with duodenal ulcer had an average antibody titer significantly higher than all the other diseases. Altogether, the ELISA was found to be able to predict 75.3% of the patients with any gastroduodenal disease and 100% of the patients with duodenal ulcer.

In one particular ELISA, a recombinant protein containing 230 amino acids deriving from CAI antigen was identified by screening an expression library of H. pylori DNA using an antiserum specific for the protein. The recombinant antigen was expressed as a fusion protein in E. coli, purified to homogeneity, and used to coat microtiter plates. The plates were then incubated for 90 minutes with 1/2000 of dilution goat anti-human IgG phosphatase cojugate. Following washing, the enzyme substrate was added to the plates and the optical density at 405 nm was read 30 minutes later. The cutoff level was determined by the mean absorbance plus two standard deviations, using sera from 20 individuals that had neither gastric disease nor detectable anti-H. pylori antibodies in The ELISA assay was tested on the Western blotting. peripheral blood samples of eighty-two dyspeptic patients (mean age 50.6±13.4 years, ranging from 28 to 80) undergoing routine upper gastrointestinal endoscopy examination. gastric antral mucosa of patients was obtained for histology Twenty of the patients had duodenal and Giemsa strain. ulcer, 5 had gastric ulcer, 43 had chronic active gastritis type B, 8 had duodenitis and 6 had a normal histology of gastric mucosa. All of the patients with duodenal ulcer had

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an optical density value above the cutoff level. The patients with duodenitis, gastric ulcer, and chronic gastritis, had a positive ELISA value in 75%, 80% and 53.9% of the cases, respectively. The agreement between ELISA and histological Giemsa staining was 95% in duodenal ulcer, 98% in duodenitis, 80% in gastric ulcer and 55.8% in chronic gastritis. This assay gives an excellent correlation with duodenal ulcer disease (p. 0.0005).

iii. Heat shock protein (hsp)

10 1. Materials and methods

a. <u>H. pylori</u> strains and growth conditions

H. pylori strains used were: CCUG 17874, G39 and G33 (isolated from gastric biopsies in the hospital of Grosseto, Italy), Pylo 2U+ and Pylo 2U- (provided by F. Megraud, hospital Pellegzin, Bordeaux, France), BA96 (isolated by gastric biopsies at the University of Siena, Italy). Strain Pylo 2U+ is noncytotoxic; strain Pylo 2U- is noncytotoxic and urease-negative. All strains were routinely grown on Columbia agar containing 0.2% of cyclodextrin, 5µg/ml of cefsulodin 5µg/ml and amphotericin B under microaerophilic conditions for 5-6 days at 37°C°. Cells were harvested and washed with PBS. The pellets were resuspended in Laemmli sample buffer and lysed by boiling.

Sera of patients affected by gastritis and ulcers (provided by A. Ponzetto, hospital "Le Molinette", Torino, Italy) and sera of patients with gastric carcinoma (provided by F. Roviello, University of Siena, Italy) were used.

b. Immunoscreening of the library

Five hundred thousand plaques of a \(\lambda\)gt11 <u>H. pylori</u>

DNA expression library were mixed with 5 ml of a suspension of <u>E. coli</u> strain Y1090 grown O/N in LB with 0.2% Maltose and 10mM MgSO₄, and resuspended in 10mM MgSO₄ at 0.5 O.D. After 10 minutes incubation at 37°C, 75 ml of melted TopAgarose were poured in the bacterial/phage mix and the whole was plated on BBL plates (50,000 plaques/plate). After 3.5 hrs incubation of the plated library at 42°C, nitrocellulose filters (Schleicher and Schuell, Dassel, Germany), previously wet with 10mM IPTG, were set on plates

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and incubation was prolonged for 3.5 hrs at 37°C and then O/N at 4°C. Lifted filters with lambda proteins were rinse in PBS, and saturated in 5% nonfat dried milk dissolved in TBST (10mM TRIS pH 8, 100mM NaCl, 5M MgCl₂) for 20'. The first hybridization step was performed with the sera of patients; to develop and visualize positive plaques we used an anti human Ig antibody alkaline phosphatase conjugated (Cappel, West Chester, PA) and the NBT/BCIP kit (Promega, Madison, WI) in AP buffer (100mM Tris pH 9.5, 100mM NaCl, 5mM MgCl₂) according to the manufacturer instructions.

c. Recombinant DNA procedures

Reagents and restriction enzymes used were from Sigma (St. Louis, MO) and Boehringer (Mannheim, Germany). Standard techniques were used for molecular cloning, single-stranded DNA purification, transformation in <u>E. coli</u>, radioactive labeling of probes, colony screening of the <u>H. pylori</u> DNA genomic library, Southern blot analysis, PAGE and Western blot analysis.

d. DNA sequence analysis

The DNA fragments were subcloned in Bluescript SK+
(Stratagene, San Diego, CA). Single-stranded DNA sequencing
was performed by using [33P]adATP (New England Nuclear,
Boston, MA) and the Sequenase kit (U.S. Biochemical Corp.,
Cleveland, OH) according to the manufacturer instructions.

The sequence was determined in both strands and each strand
was sequenced, on average, twice. Computer sequence
analysis was performed using the GCG package.

e. Recombinant proteins

MS2 polymerase fusion proteins were produced using the vector pEX34A, a derivative of pEX31. Insert Hp67 (from nucleotide 445 to nucleotide 1402 in Fig. 5), and the EcoRI linkers were cloned in frame into the EcoRi site of the vector. In order to confirm the location of the stop codon, the HpG3' HindIII fragment was cloned in frame into the HindIII site of pEX34A. Recombinant plasmids were transformed in E. coli K12 [H1 Atrp. In both cases after induction, a fusion protein of the expected molecular weight was produced. In the case of the EcoRI/EcoRI fragment, the fusion protein obtain after induction was electroeluted to

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immunize rabbits using standard protocols.

2. Results

a. Screening of an expression library and cloning of <u>H.</u> pylori hsp

In order to find a serum suitable for the screening of an <u>H. pylori</u> DNA expression library, sonicated extracts of <u>H. pylori</u> strain CCUG 17874 were tested in Western blot analysis against sera of patients affected by different forms of gastritis. The pattern of antigen recognition by different sera was variable, probably due to differences in the individual immune response as well as to the differences in the antigens expressed by the strains involved in the infection.

Serum N°19 was selected to screen a lgt11 H. <u>pylori</u> DNA expression library to identify <u>H. pylori</u> specific 15 antigens, expressed in vivo during bacterial growth. Following screening of the library with this serum, many positive clones were isolated and characterized. The nucleotide sequence of one of these, called Hp67, revealed an open-reading frame of 958 base-pairs, coding for a 20 protein with high homology to the hsp60 family of heat-shock proteins, Ellis, Nature 358:191-92 (1992). In order to obtain the entire coding region, we used fragment Hp67 as a probe on Southern blot analysis of H. pylori DNA digested with different restriction enzymes. Probe Hp67 recognized 25 two HindIII bands of approximately 800 and 1000 base-pairs, respectively. A genomic <u>H. pylori</u> library of HindIIIdigested DNA was screened with probe Hp67 and two positive clones (HpG5' and HpG3') of the expected molecular weight 30 were obtained. E. coli containing plasmids pHp60G2 (approximately nucleotides 1 to 829) and (approximately nucleotides 824 to 1838) were deposited with the American Type Culture Collection (ATCC).

b. Sequence analysis

The nucleotide sequence analysis revealed an openreading frame of 1638 base-pairs, with a putative ribosome binding site 6 base-pairs upstream the starting ATG. Fig. 5 shows the nucleotide and amino acid sequences of <u>H. pylori</u> hsp. The putative ribosome-binding and the internal HindIII

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site are underlined. Cytosine in position 445 and guanine in position 1402 are the first and last nucleotide, respectively, in fragment Hp67. Thymine 1772 was identified as the last putative nucleotide transcribed using an algorithm for the localization of factor-independent terminator regions. The open-reading frame encoded for a protein of 546 amino acids, with a predicted molecular weight of 58.3 KDa and a predicted pI of 5.37. The codon preference of this gene is in agreement with the <u>H. pylori</u> codon usage.

The analysis of the hydrophylicity profiles revealed a protein mostly hydrophilic, without a predicted leader peptide or other transmembrane domains. The amino terminal sequence showed 100% homology to the sequence of 30 amino acids determined by Dunn et al., Infect. Immun. 60:1946-51 (1992) on the purified protein and differed by only on reside (Ser42 instead of Lys) from the sequence of 44 amino acids published by Evans et al, Infect. Immun. 60:2125-27 (1992). (Evans et al., 1992). The N-terminal sequence of the mature hsp protein did not contain the starting methionine, indicating that this had been removed after translation.

c. Homology with hsp60 family

The amino acid sequence analysis showed a very strong homology with the family of heat-shock proteins hsp60, whose members are present in every living organism. Based on the degree of homology between hsp60 proteins of different species, <u>H. pylori</u> hsp belongs to the subgroup of hsp60 proteins of Gram negative bacteria; however, the degree of homology to the other proteins of the hsp60 family is very high (at least 54% identity).

d. Expression of recombinant proteins and production of a polyclonal antiserum

The inserts of clone Hp67 and of clone HpG3' were subcloned in the expression vector pEX34A in order to express these open-reading frames fused to the aminoterminus of the MS2 polymerase. The clones produced recombinant proteins of the expected size and were recognized by the human serum used for the initial screening. The fused

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protein derived from clone Hp67 was electroeluted and used to immunize rabbits in order to obtain anti-hsp specific polyclonal antisera. The antiserum obtained recognized both fusion proteins, and a protein of 58 KDa on whole-cell extracts of several strains of <u>H. pylori</u> tested, including a urease-negative strain and noncytotoxic strains.

Hsp has been shown to be expressed by all the H. <u>pylori</u> strains tested and its expression is not associated with the presence of the urease or with the cytotoxicity. The protein recognized by the anti-hsp antiserum was found in the water soluble extracts of H. pylori and copurified with the urease subunits. This suggests a weak association of this protein with the outer bacterial membrane. hsp can be described as urease-associated and surface exposed. The cellular surface localization is surprising as most of the hsp homologous proteins are localized in the cytoplasm or in mitochondria and plastids. The absence of a leader peptide in hsp suggests that this is either exported to the membrane by a peculiar export system, or that the protein is released from the cytoplasm and is passively adsorbed by the bacterial membrane after death of the bacterium.

Hsp60 proteins have been shown to act as molecular chaperons assisting the correct folding, assembly and translocation of either oligomeric or multimeric proteins. The cellular localization of H. pylori hsp and its weak association with urease suggest that hsp may play a role in assisting the folding and/or assembly of proteins exposed on the membrane surface and composed of multiple subunits such as the urease, whose final quaternary structure is ABB. Austin et al., J. Bacteriol. 174:7470-73 (1992) showed that the <u>H. pylori</u> hsp ultrastructure is composed of seven subunits assembled in a disk-shaped particle that further stack side by side in groups of four. This structure resembles the shape dimension and of the macromolecule and this could explain the common properties these macromolecules of two that lead to their copurification. H. pylori hsp gene, however, is not part of the urease operon. In agreement with the gene structure of

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other bacterial hsp60 proteins, it should be part of a dicistronic operon.

e. Presence of anti-hsp antibodies in patients with gastroduodenal diseases

The purified fusion protein was tested by Western blot using sera of patients infected by H. pylori and affected by atrophic and superficial gastritis, and patients with duodenal and gastric ulcers: most of the sera recognized the recombinant protein. However, the degree of recognition greatly varied between different individuals and . the antibody levels did not show any obvious correlation with the type of disease. In addition, antibodies against H. pylori antigens and in particular against hsp protein were found in most of the 12 sera of patients affected by gastric carcinoma that were tested. Although H. pylori hsp recognition could not be put in relation with a particular clinical state of the disease given the high conservation between <u>H. pylori</u> hsp and its human homolog, it is possible that this protein may induce autoimmune antibodies crossreacting with the human counterpart. This class of homologous proteins has been implicated in the induction of autoimmune disorders in different systems. Thenpresent of high titers of anti-H. pylor hsp antibodies, potentially cross-reacting with the human homolog in dispeptic patients, suggests that this protein has a role in gastroduodenal This autoreactivity could play a role in the disease. tissue damage that occurs in H. pylori-induced gastritis, thus increasing the pathogenic mechanisms involved in the infection of this bacterium.

The high levels of antibodies against such a conserved protein is somewhat unusual; due to the high homology between members of the hsp60 family, including the human one, this protein should be very well tolerated by the host immune system. The strong immune response observed in many patients may be explained in two different ways: (1) the immune response is directed only against epitopes specific for <u>H. pylori</u> hsp; (2) the immune response is directed against epitopes which are in common between <u>H. pylori</u> hsp and human homolog.

H. Deposit of Biological Materials

The following materials were deposited on December 15, 1992 and January 22, 1993 by Biocine Sclavo, S.p.A., the assignee of the present invention, with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland, phone (301) 231-5519, under the terms of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for Purposes of Patent Procedure. For the cytotoxin protein (CT):

ATCC No. 69157 <u>E. coli</u> TG1 containing the plasmid TOXHH1 ATCC No. n/a <u>E. coli</u> TG1 containing the plasmid TOXEE1 For the CAI protein:

ATCC No. 69158 E. coli TG1 containing the plasmid 57/D

ATCC No. 69159 E. coli TG1 containing the plasmid 64/4

ATCC No. 69160 <u>E. coli</u> TG1 containing the plasmid P1-24 ATCC No. 69161 <u>E. coli</u> TG1 containing the plasmid B/1 For the heat shock protein (hsp):

ATCC No. 69155 <u>E. coli</u> TG1 containing the plasmid pHp60G2 ATCC No. 69156 <u>E. coli</u> TG1 containing the plasmid pHp605

20 These deposits are provided as convenience to those of skill in the art, and are not an admission that a deposit is required under 35 U.S.C. §112. The nucleic acid sequences of these deposits, as well as the amino acid sequences the polypeptides of encoded thereby, are incorporated herein by reference and should be referred to 25 in the event of any error in the sequences described herein as compared with the sequences of the deposits. A license may be required to make, use, or sell the deposited materials, and no such license is granted hereby.

CLAIMS

What is claimed is:

A recombinant <u>Helicobacter pylori</u> protein, or
 a derivative or fragment thereof.

2. The recombinant protein according to claim 1 wherein the protein is a <u>Helicobacter pylori</u> cytotoxin or a precursor, derivative or fragment thereof.

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3. The recombinant protein according to claim 2 wherein the cytotoxin, precursor, derivative or fragment thereof has the amino acid sequence of Figure 2, or a portion thereof.

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4. The recombinant protein according to claim 1 wherein the protein is a <u>Helicobacter pylori</u> cytotoxin associated immunodominant antigen, or a derivative or fragment thereof.

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5. The recombinant protein according to claim 4 wherein the cytotoxin associated immunodominant antigen, derivative or fragment has the amino acid sequence of Figure 4, or a portion thereof.

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- 6. The recombinant protein according to claim 1 wherein the protein is a <u>Helicobacter pylori</u> heat shock protein, or a derivative or fragment thereof.
- 7. The recombinant protein according to claim 6, wherein the heat shock protein, derivative or fragment has the amino acid sequence of Figure 5 or a portion thereof.
- 8. The recombinant protein according to claim 2 or 3 wherein the recombinant protein exhibits substantially no toxicity, or substantially reduced toxicity.
 - 9. The recombinant protein according to any one of claims 4 to 7 wherein the recombinant protein is

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immunogenic and exhibits no functional contribution to toxicity, or a substantially reduced functional contribution to toxicity.

- or 9 wherein the recombinant protein is chemically modified to reduce or abolish toxicity or functional contribution to toxicity.
- 11. The recombinant protein according to claim 8 or 9 wherein the recombinant protein contains one or more amino acid substitutions or deletions.
- 12. The recombinant protein according to any one of the preceding claims which is labelled or coupled to a solid support.
- of claims 1 to 11 for use in the treatment of <u>Helicobacter</u> pylori infection.
 - 14. The recombinant protein according to any one of claims 1 to 11 for use as a vaccine.
- 15. A vaccine or therapeutic composition comprising a recombinant protein according to any one of claims 1 to 11 and a pharmaceutically acceptable carrier.
- 16. The vaccine or therapeutic composition according to claim 15 comprising two or more recombinant proteins according to any one claims 1 to 11.
- 17. The vaccine or therapeutic composition according to claim 16 comprising, in combination, two or more of
 - i) a recombinant <u>Helicobacter pylori</u> cytotoxic protein precursor, derivative or fragment thereof,
 - ii) a <u>Helicobacter pylori</u> recombinant cytotoxin

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associated immunodominant antigen, or a derivative or fragment thereof,

- iii) <u>Helicobacter pylori</u> recombinant heat shock protein or a derivative or fragment thereof and/or
 - iv) a <u>Helicobacter pylori</u> urease.
- 18. The vaccine or therapeutic composition according to any one of claims 15 to 17 comprising an adjuvant.

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- 19. A method for the preparation of a vaccine or therapeutic composition according to any one of claims 15 or 18 comprising bringing one or more recombinant proteins according to any one of claims 1 to 11 into association with a pharmaceutically acceptable carrier and optionally an adjuvant.
- 20. An immunodiagnostic assay comprising at least one step involving as at least one binding partner, a recombinant protein according to any one of claims 1 to 12, optionally labelled or coupled to a solid support.
- 21. An immunodiagnosis kit for performing an assay according to claim 20, comprising at least one recombinant protein according to any one of claims 1 to 20.
 - 22. Use of one or more recombinant proteins according to any one of claims 1 to 11 for the manufacture of a medicament for the treatment of <u>Helicobacter pylori</u> infection.
 - 23. A method of treatment of an individual infected with <u>Helicobacter pylori</u> comprising administering an effective amount of a recombinant protein according to 1 to 11.
 - 24. The method of treatment according to claim 23 comprising administering an effective amount of, in combination, two or more of

- i) a recombinant <u>Helicobacter pylori</u> cytotoxic protein precursor, derivative or fragment thereof,
- ii) a <u>Helicobacter pylori</u> recombinant cytotoxin associated immunodominant antigen, or a derivative or fragment thereof,
- iii) a <u>Helicobacter pylori</u> recombinant heat shock protein or a derivative or fragment thereof and/or
 - iv) a <u>Helicobacter pylori</u> urease.
- 25. A method of vaccination comprising administering an immunologically effective amount of, in combination, two or more of
 - i) a recombinant <u>Helicobacter pylori</u> cytotoxic protein precursor, derivative or fragment thereof,

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- ii) a <u>Helicobacter pylori</u> recombinant cytotoxin associated immunodominant antigen, or a derivative or fragment thereof,
- iii) a <u>Helicobacter pylori</u> recombinant heat shock 20 protein or a derivative or fragment thereof and/or
 - iv) a <u>Helicobacter pylori</u> urease.
 - 26. A recombinant polynucleotide encoding a recombinant protein according to any one of claims 1 to 11.

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27. A recombinant polynucleotide encoding a <u>Helicobacter pylori</u> cytotoxic protein or a derivative or fragment thereof comprising all or part of the nucleotide sequence of Figure 1.

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- 28. A recombinant polynucleotide encoding a <u>Helicobacter pylori</u> recombinant cytotoxin associated immunodominant antigen or a derivative or fragment thereof comprising all or a part of the nucleotide sequence of Figure 4.
- 29. A recombinant polynucleotide encoding a <u>Helicobacter pylori</u> recombinant heat shock protein or a derivative or fragment thereof comprising all or a part of

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the nucleotide sequence of Figure 5.

- 30. A polynucleotide probe comprising all or part of the recombinant polynucleotide according to any one of claims 26 to 29.
- 31. A nucleic acid assay wherein in at least one step involves a polynucleotide probe according to claim 30.
- 32. A kit for performing a nucleic acid assay comprising at least one polynucleotide probe according to claim 30.
- 33. A polynucleotide amplification process employing a polynucleotide primer wherein in at least one primer is a recombinant polynucleotide comprising all or part of the recombinant polynucleotide according to any one of claims 26 to 29.
- 34. A kit for performing a polynucleotide amplification process employing a polynucleotide primer wherein in at least one primer is a recombinant polynucleotide comprising all or part of the recombinant polynucleotide according to any one of claims 26 to 29.
 - 35. A vector comprising a recombinant polynucleotide according to any one of claims 26 to 29.
- 36. A host cell transformed with a vector according to claim 35.
 - 37. A method for the production of a recombinant polypeptide according to any one of claims 1 to 11, comprising culturing a host cell according to claim 36 and isolating the recombinant polypeptide.

1/14

1	AAAAAGAAAG	GAAGAAAATG	GAAATACAAC	AAACACACCG	CAAAATCAAT
51	CGCCCTCTGG	TTTCTCTCGC	TTTAGTAGGA	GCATTAGTCA	GCATCACACC
101	GCAACAAAGT	CATGCCGCCT	TTTTCACAAC	CGTGATCATT	CCAGCCATTG
151	TTGGGGGTAT	CGCTACAGGC	ACCGCTGTAG	GAACGGTCTC	AGGGCTTCTT
201	AGCTGGGGGC	TCAAACAAGC	CGAAGAAGCC	AATAAAACCC	CAGATAAACC
251	CGATAAAGTT	TGGCGCATTC	AAGCAGGAAA	AGGCTTTAAT	GAATTCCCTA
301	ACAAGGAATA	CGACTTATAC	AGATCCCTTT	TATCCAGTAA	GATTGATGGA
351	GGTTGGGATT	GGGGGAATGC	CGCTAGGCAT	TATTGGGTCA	AAGGCGGGCA
401	ACAGAATAAG	CTTGAAGTGG	ATATGAAAGA	CGCTGTAGGG	ACTTATACCT
451	TATCAGGGCT	TAGAAACTTT	ACTGGTGGGG	ATTTAGATGT	CAATATGCAA
501	AAAGCCACTT	TACGCTTGGG	CCAATTCAAT	GGCAATTCTT	TTACAAGCTA
551	TAAGGATAGT	GCTGATCGCA	CCACGAGAGT	GATTTCAACG	CTAAAAATAT
601	CTCAATTGAT	AATTTTGCAG	AAATCAACAA	CTCGTGTGGG	TTCTGGAGCC
651	GGGAGGAAAG	CCAGCTCTAC	GGTTTTGACT	TTGCAAGCTT	CAGAAGGGAT
701	CACTAGCGAT	AAAAACGCTG	AAATTTCTCT	TTATGATGGT	GCCACGCTCA
751	ATTTGGCTTC	AAGCAGCGTT	AAATTAATGG	GTAATGTGTG	GATGGGCCGT
801	TTGCAATACG	TGGGAGCGTA	TTTGGCCCCT	TCATACAGCA	CGĂTAAACAC
851	TTCAAAAGTA	ACAGGGGAAG	TGAATTTTAA	CCACCTCACT	GTTGGCGATA
901	AAAACGCCGC	TCAAGCGGGC	ATTATCGCTA	ATAAAAAGAC	TAATATTGGC
951	ACACTGGATT	TGTGGCAAAG	CGCCGGGTTA	AACATTATCG	CTCCTCCAGA
1001	AGGTGGCTAT	AAGGATAAAC	CCAATAATAC	CCCTTCTCAA	AGTGGTGCTA
1051	AAAACGACAA	AAATGAAAGC	GCTAAAAACG	ACAAACAAGA	GAGCAGTCAA
1101	AATAATAGTA	ACACTCAGGT	CATTAACCCA	CCCAATAGTG	CGCAAAAAAC
1151	AGAAGTTCAA	CCCACGCAAG	TCATTGATGG	GCCTTTTGCG	GGCGGCAAAG
1201	ACACGGTTGT	CAATATCAAC	CGCATCAACA	CTAACGCTGA	TGGCACGATT
1251	AGAGTGGGAG	GGTTTAAAGC	TTCTCTTACC	ACCAATGCGG	CTCATTTGCA
1301	TATCGGCAAA	GGCGGTGTCA		TCAAGCGAGC	GGGCGCTCTC
FIG. 1A					

2/14 -

1351 TTATAGTGGA AAATCTAACT GGGAATATCA CCGTTGATGG GCCTTTAAGA 1401 GTGAATAATC AAGTGGGTGG CTATGCTTTG GCAGGATCAA GCGCGAATTT 1451 TGAGTTTAAG GCTGGTACGG ATACCAAAAA CGGCACAGCC ACTTTTAATA 1501 ACGATATTAG TCTGGGAAGA TTTGTGAATT TAAAGGTGGA TGCTCATACA 1551 GCTAATTTTA AAGGTATTGA TACGGGTAAT GGTGGTTTCA ACACCTTAGA 1601 TTTTAGTGGC GTTACAGACA AAGTCAATAT CAACAAGCTC ATTACGGCTT 1651 CCACTAATGT GGCCGTTAAA AACTTCAACA TTAATGAATT GATTGTTAAA 1701 ACCAATGGGA TAAGTGTGGG GGAATATACT CATTTTAGCG AAGATATAGG 1751 CAGTCAATCG CGCATCAATA CCGTGCGTTT GGAAACTGGC ACTAGGTCAC 1801 TTTTCTCTGG GGGTGTTAAA TTTAAAGGTG GCGAAAAATT GGTTATAGAT 1851 GAGTTTTACT ATAGCCCTTG GAATTATTTT GACGCTAGAA ATATTAAAAA 1901 TGTTGAAATC ACCAATAAAC TTGCTTTTGG ACCTCAAGGA AGTCCTTGGG 1951 GCACATCAAA ACTTATGTTC AATAATCTAA CCCTAGGTCA AAATGCGGTC 2001 ATGGATTATA GCCAATTTTT AAATTTAACC ATTCAAGGGG ATTTCATCAA 2051 CAATCAAGGC ACTATCAACT ATCTGGTCCG AGGTGGGAAA GTGGCAACCT 2101 TAAGCGTAGG CAATGCAGCA GCTATGATGT TTAATAATGA TATAGACAGC 2151 GCGACCGGAT TTTACAAACC GCTCATCAAG ATTAACAGCG CFCAAGATCT 2201 CATTAAAAAT ACAGAACATG TTTTATTGAA AGCGAAAATC ATTGGTTATG 2251 GTAATGTTTC TACAGGTACC AATGGCATTA GTAATGTTAA TCTAGAAGAG 2301 CAATTCAAAG AGCGCCTAGC CCTTTATAAC AACAATAACC GCATGGATAC 2351 TTGTGTGGTG CGAAATACTG ATGACATTAA AGCATGCGGT ATGGCTATCG 2401 GCGATCAAAG CATGGTGAAC AACCCTGACA ATTACAAGTA TCTTATCGGT 2451 AAGGCATGGA AAAATATAGG GATCAGCAAA ACAGCTAATG GCTCTAAAAT 2501 TTCGGTGTAT TATTTAGGCA ATTCTACGCC TACTGAGAAT GGTGGCAATA 2551 CCACAAATTT ACCCACAAAC ACCACTAGCA ATGCACGTTC TGCCAACAAC 2601 GCCCTTGCAC AAAACGCTCC TTTCGCTCAA CCTAGTGCTA CTCCTAATTT 2651 AGTCGCTATC AATCAGCATG ATTTTGGCAC TATTGAAAGC GTGTTTGAAT

FIG. 1B

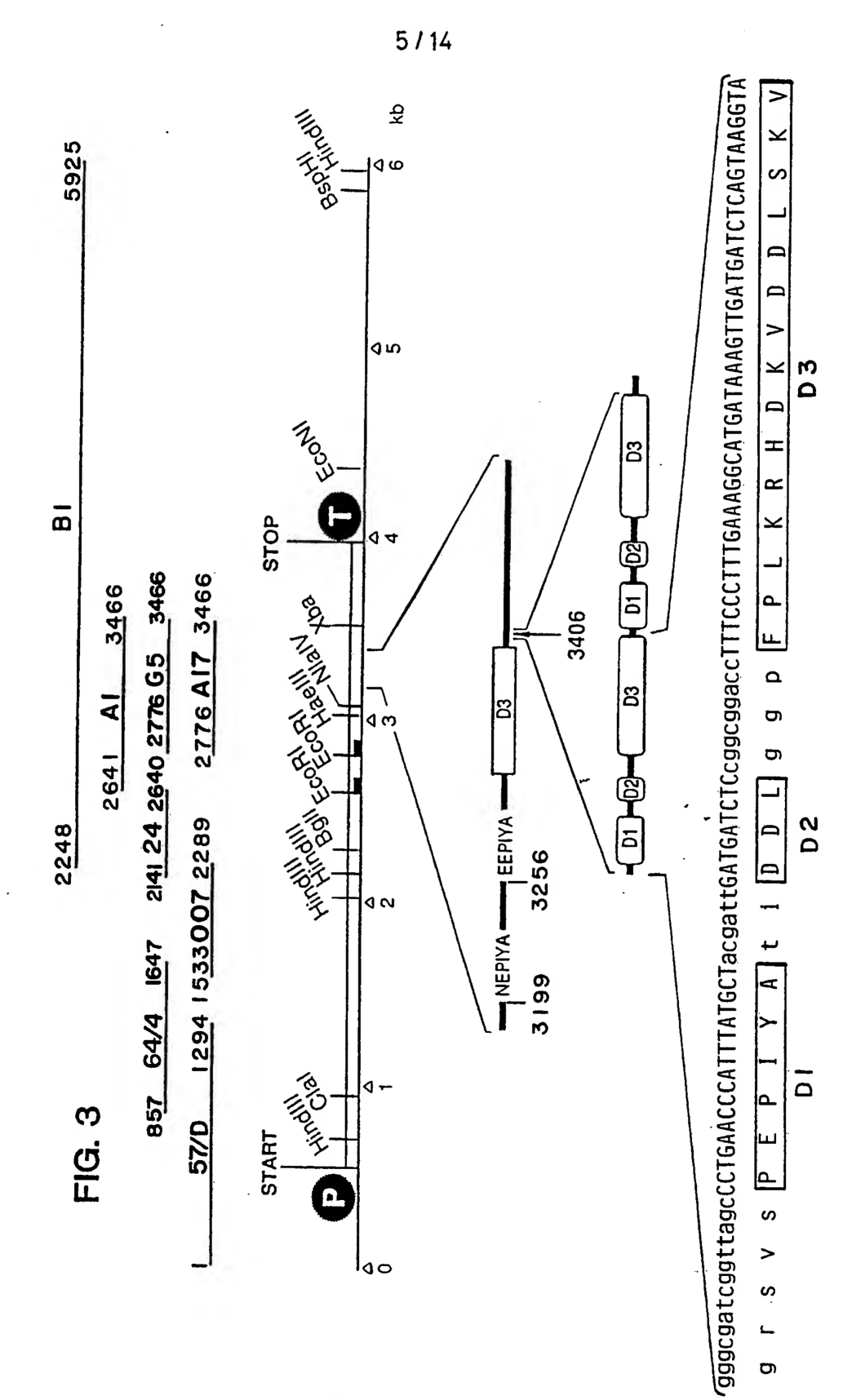
3/14

2701 TGGCTAACCG CTCTAAAGAT ATTGACACGC TTTATGCTAA CTCAGGCGCT 2751 CAAGGCAGGG ATCTCTTACA AACCTTATTG ATTGATAGCC ATGATGCGGG 2801 TTATGCCAGA AAAATGATTG ATGCTACAAG CGCTAATGAA ATCACCAAGC 2851 AATTGAATAC GGCCACTACC ACTTTAAACA ACATAGCCAG TTTAGAGCAT 2901 AAAACCAGCG GCTTACAAAC TTTGAGCTTG AGTAATGCGA TGATTTTAAA 2951 TTCTCGTTTA GTCAATCTCT CCAGGAGACA CACCAACCAT ATTGACTCGT 3001 TCGCCAAACG CTTACAAGCT TTAAAAGACC AAAAATTCGC TTCTTTAGAA 3051 AGCGCGGCAG AAGTGTTGTA TCAATTTGCC CCTAAATATG AAAAACCTAC 3101 CAATGTTTGG GCTAACGCTA TTGGGGGAAC GAGCTTGAAT AATGGCTCTA 3151 ACGCTTCATT GTATGGCACA AGCGCGGGCG TAGACGCTTA CCTTAACGGG 3201 CAAGTGGAAG CCATTGTGGG CGGTTTTGGA AGCTATGGTT ATAGCTCTTT 3251 TAATAATCGT GCGAACTCCC TTAACTCTGG GGCCAATAAC ACTAATTTTG 3301 GCGTGTATAG CCGTATTTTA ACCAACCAGC ATGAATTTGA CTTTGAAGCT 3351 CAAGGGCAC TAGGGAGCGA TCAATCAAGC TTGAATTTCA AAAGCGCTCT 3401 ATTACAAGAT TTGAATCAAA GCTATCATTA CTTAGCCTAT AGCGCTGCAA 3451 CAAGAGCGAG CTATGGTTAT GACTTCGCGT TTTTTAGGAA CGCTTTAGTG 3501 TTAAAACCAA GCGTGGGTGT GAGCTATAAC CATTTAGGTT CAACCAACTT 3551 TAAAAGCAAC AGCACCAATC AAGTGGCTTT GAAAAATGGC TCTAGCAGTC 3601 AGCATTTATT CAACGCTAGC GCTAATGTGG AAGCGCGCTA TTATTATGGG 3651 GACACTTCAT ACTTCTACAT GAATGCTGGA GTTTTACAAG AGTTCGCTCA 3701 TGTTGGCTCT AATAACGCCG CGTCTTTAAA CACCTTTAAA GTGAATGCCG 3751 CTCGCAACCC TTTAAATACC CATGCCAGAG TGATGATGGG TGGGGAATTA 3801 AAATTAGCTA AAGAAGTGTT TTTGAATTTG GGCGTTGTTT ATTTGCACAA 3851 TTTGATTTCC AATATAGGCC ATTTCGCTTC CAATTTAGGA ATGAGGTATA 3901 GTTTCTAAAT ACCGCTCTTA AACCCATGCT CAAAGCATGG GTTTGAAATC 3951 TTACAAAACA

FIG. 1C

1 MEIQQTHRKI NRPLVSLALV GALVSITPQQ SHAAFFTTVI IPAIVGGIAT 51 GTAVGTVSGL LSWGLKQAEE ANKTPDKPDK VWRIQAGKGF NEFPNKEYDL 101 YRSLLSSKID GGWDWGNAAR HYWVKGGQQN KLEVDMKDAV GTYTLSGLRN 151 FTGGDLDVNM QKATLRLGQF NGNSFTSYKD SADRTTRVIS TLKISQLIIL 201 QKSTTRVGSG AGRKASSTVL TLQASEGITS DKNAEISLYD GATLNLASSS 251 VKLMGNVWMG RLQYVGAYLA PSYSTINTSK VTGEVNFNHL TVGDKNAAQA 301 GIIANKKTNI GTLDLWQSAG LNIIAPPEGG YKDKPNNTPS QSGAKNDKNE 351 SAKNDKQESS QNNSNTQVIN PPNSAQKTEV QPTQVIDGPF AGGKDTVVNI 401 NRINTNADGT IRVGGFKASL TTNAAHLHIG KGGVNLSNQA SGRSLIVENL 451 TGNITVDGPL RVNNQVGGYA LAGSSANFEF KAGTDTKNGT ATFNNDISLG 501 RFVNLKVDAH TANFKGIDTG NGGFNTLDFS GVTDKVNINK LITASTNVAV 551 KNFNINELIV KTNGISVGEY THFSEDIGSQ SRINTVRLET GTRSLFSGGV 601 KFKGGEKLVI DEFYYSPWNY FDARNIKNVE ITNKLAFGPQ GSPWGTSKLM 651 FNNLTLGQNA VMDYSQFLNL TIQGDFINNQ GTINYLVRGG KVATLSVGNA 701 AAMMENNDID SATGEYKPLI KINSAQDLIK NTEHVLLKAK IIGYGNVSTG 751 TNGISNVNLE EQFKERLALY NNNNRMDTCV VRNTDDIKAC GMAIGDQSMV 801 NNPDNYKYLI GKAWKNIGIS KTANGSKISV YYLGNSTPTE NGGNTTNLPT 851 NTTSNARSAN NALAQNAPFA QPSATPNLVA INQHDFGTIE SVFELANRSK 901 DIDTLYANSG AQGRDLLQTL LIDSHDAGYA RKMIDATSAN EITKQLNTAT 951 TTLNNIASLE HKTSGLQTLS LSNAMILNSR LVNLSRRHTN HIDSFAKRLQ 1001 ALKDQKFASL ESAAEVLYQF APKYEKPTNV WANAIGGTSL NNGSNASLYG 1051 TSAGVDAYLN GQVEAIVGGF GSYGYSSFNN RANSLNSGAN NTNFGVYSRI 1101 LTNQHEFDFE AQGALGSDQS SLNFKSALLQ DLNQSYHYLA YSAATRASYG 1151 YDFAFFRNAL VLKPSVGVSY NHLGSTNFKS NSTNQVALKN GSSSQHLFNA 1201 SANVEARYYY GDTSYFYMNA GVLQEFAHVG SNNAASLNTF KVNAARNPLN 1251 THARVMMGGE LKLAKEVFLN LGVVYLHNLI SNIGHFASNL GMRYSF

FIG. 2



ERSATZBLATT

4

AATAATCTTCAAGTAGCTTTTCTTAAAGTTGATAACGCTGTCGCTTCATACGATCCTGAT

23 N N L Q V A F L K V D N A V A S Y D P D

CAATTAAGGGAAGAATACTCCAATAAAGCGATCAAAAAATCCTACCAAAAAGAATCAGTAT

63 Q L R E E Y S N K A I K N P T K K N Q Y
GAATCTTCCACAAAGAGCTTTCAGAAATTTGGGGATCAGCGTTACCGAATTTTCACAAGT

103 E S S T K S F Q K F G D Q R Y R I F T S GAAAATATCATACAACCCCCTATCCTTGATGATAAAGAGAAAGCGGAGTTTTTGAAATCT

143 E N I I Q P P I L D D K E K A E F L K S
ATGGGCGTGTTTGATGAGTCCTTGAAAGAAGGCAAGAAGCAGAAAAAAATGGAGAGCCT

183 M G V F D E S L K E R Q E A E K N G E P
GATGTCAAAGAAGCAATCAATCAAGAACCAGTTCCCCATGTCCAACCAGATATAGCCACT

223 D V K E A I N Q E P V P H V Q P D I A T AATTTTTCTAAATTCACTCTTGGCGATATGGAAATGTTAGATGTTGAGGGAGTCGCTGAC

263 N F S K F T L G D M E M L D V E G V A D
TTAATGGGGAGTCATAATGGCATAGAACCTGAAAAAGTTTCATTGTTGTATGGGGGCAAT

303 L M G S H N G I E P E K V S L L Y G G N AACAATGTGGCTACAATAATTAATGTGCATATGAAAAACGGCAGTGGCTTAGTCATAGCA

343 N N V A T I I N V H M K N G S G L V I A GGCTCACAACGAGCATTAAGTCAAGAAGAGAGATCCAAAACAAAATAGATTTCATGGAATTT

383 G S Q R A L S Q E E I Q N K I D F M E F ACTGAGATTAAAGATTTCCAAAAAGACTCTAAGGCTTATTTAGACGCCCTAGGGAATGAT

423 T E I K D F Q K D S K A Y L D A L G N D AATGGGGATTTGAGGTTTAGAT

463 N G D L S Y T L K D Y G K K A D K A L D
TATTCTAATTCAAATACACCAACGCCTCCAAGAATCCCAATAAGGGTGTAGGCGTTACG

FIG. 4A

ALCIGICCIALIGATITGTTTTCCATTTTGTTTCCCATGTGGATCTTGTGGATCACAAAC	120
CATGTGCTCACCTTGACTAACCATTTCTCCAACCATACTTTAGCGTTGCATTTGATTTCT	240
TTGTTAATTGTGGGTAAAAATGTGAATCGTCCTAGCCTTTAGACGCCTGCAACGATCGGG	360
AATGAGAATGTTCAAAGACATGAATTGACTACTCAAGCGTGTAGCGATTTTTAGCAGTCT	480
`AACGAAACCATTGACCAACAACCACAAACCGAAGCGGCTTTTAACCCGCAGCAATTTATC	600
NETIDQQPQTEAAFNPQQFI	
CAAAAACCAATCGTTGATAAGAACGATAGGGATAACAGGCAAGCTTTTGAAGGAATCTCG	720
QKPIVDKNDRDNRQAFEGIS	
TTTTCAGACTTTATCAATAAGAGCAATGATTTAATCAACAAAGACAATCTCATTGATGTA	840
F S D F I N K S N D L I N K D N L I D V	
TGGGTGTCCCATCAAAACGATCCGTCTAAAATCAACACCCGATCGAT	960
WVSHQNDPSKINTRSIRNFM	
GCCAAACAATCTTTTGCAGGAATCATTATAGGGAATCAAATCCGAACGGATCAAAAGTTC	1080
AKQSFAGIIIGNQIRTDQKF	
ACTGGTGGGGATTGGTTGGATATTTTTCTCTCATTTATATTTGACAAAAAAAA	1200
TGGDWLDIFLSFIFDKKQSS	
ACCACCACCGACATACAAGGCTTACCGCCTGAAGCTAGAGATTTACTTGATGAAAGGGGT	1320
TTTDIQGLPPEARDLLDERG	
ATTGATCCCAATTACAAGTTCAATCAATTATTGATTCACAATAACGCTCTGTCTTCTGTG	1440
IDPNYKFNQLLIHNNALSSV	·
GGTGGTCCTGGAGCTAGGCATGATTGGAACGCCACCGTTGGTTATAAAGACCAACAAGGC	1560
G G P G A R H D W N A T V G Y K D Q Q G	
GGTGGTGAGAAAGGGATTAACAACCCTAGTTTTTATCTCTACAAAGAAGACCAACTCACA	1680
GGEKGINNPSFYLYKEDQLT	
CTTGCACAAAATAATGCTAAATTAGACAACTTGAGCGAGAAAGAGAAGAGAAAAATTCCGA	1800
LAQNNAKLDNLSEKEKFR	
CGTATTGCTTTTGTTTCTAAAAAAGACACAAAACATTCAGCTTTAATTACTGAGTTTGGT	1920
RIAFVSKKDTKHSALITEFG	
AGGGAGAAAAATGTTACTCTTCAAGGTAGCCTAAAACATGATGGCGTGATGTTTGTT	2040
REKNVTLQGSLKHDGVMFVD	
AATGGCGTTTCCCATTTAGAAGTAGGCTTTAACAAGGTAGCTATCTTTAATTTGCCTGAT	2160

FIG. 4B

503 Y S N F K Y T N A S K N P N K G V G V T TTAAATAATCTCGCTATCACTAGTTTCGTAAGGCGGAATTTAGAGGATAAACTAACCACT 543 L N N V R R N L : D GAATTGGTTGGAAAAACTTTAAACTTCAATAAAGCTGTAGCTGACGCTAAAAACACAGGC NKA VADA CATTTAGAGAAAGAAGTAGAGAAAAAATTGGAGAGCAAAAAGCGGCAACAAAAATAAAATG 623 H L E K E V E K K L E S K S G N K N K M GCTAATAGAGACGCAAGAGCAATCGCTTACGCTCAGAATCTTAAAGGCATCAAAAGGGAA A QNL GAATTCAAAAATGGCAAAAATAAGGATTTCAGCAAGGCAGAAGAAAAACACTAAAAGCCCTT 703 E F K D F S K A E AATGCAGCTTTGAATGAATTCAAAAATGGCAAAAATAAGGATTTCAGCAAGGTAACGCAA 743 N A A L N E F K N N AAAGTTGATAATCTCAATCAAGCGGTATCAGTGGCTAAAGCAACGGGTGATTTCAGTAGG 783 K V D N L Q A S CAAAAAAATGAAAGTCTCAATGCTAGAAAAAAATCTGAAATATATCAATCCGTTAAGAAT 823 Q K N E S L N A YQSVKN AAAAACTTTTCGGACATCAAGAAAGAGTTGAATGCAAAACTTGGAAATTTCAATAACAAT 863 K N F S D I K K E L N A K L G N F N N N CAAGCAGCTAGCCTTGAAGAACCCATTTACGCTCAAGTTGCTAAAAAGGTAAATGCAAAA 903 Q A A S L E E P I Y A Q V A K K V N A K CCTTTGAAAAGGCATGATAAAGTTGATGATCTCAGTAAGGTAGGGCTTTCAAGGAATCAA 943 PLKRHDKVDDLSKVGLSRNQ TTTGGCAATCTAGAGCAAACGATAGACAAGCTCAAAGATTCTACAAAACACAATCCCATG 983 F G N L E Q T I D K L K D S T K H

FIG.4C

N	G	٧	S	Н		Ε.			-	• •	• •	•		_	F	N		Ρ	D	
A/	AAG	GAT	TGT	CCC	CAC	AAG	AAG(CTA	ATA	AGC	TTA	TCA	AAG	ATT	TTT	TGA	GCA	GCA	ACAAA	2280
K	G	Ļ	S	P	Q	E	A	N	K	L	I	K	D	F	L	S	S	N	K	
A <i>A</i>	\TT	ATG	ATG	AAG	TGA	AAA	AAG(CTC	AGA	AAG	ATC	TTG	AAA	AAT	CTC	TAA	GGA	AAC	GAGAG	2400
N	Y	D	E	٧	K	K	Α	Q	K	D	L	Ε	K	S	L	R	K	R	Ε	
G <i>P</i>	IAG	CAA	AAG	CTC	٩AG	CTA	ACAG	SCC/	AAA	AGA	ATG	AGAT	TTT	TTG	CGT	TGA	TCA	ATA	AAGAG	2520
E	A	K	A	Q	Α	N	S	Q	K	D	Ε	I	F	Ä	L	1	N	K	Ε	
П	GT(CTGA	\TA/	ACT	ΓTG	AAA	TGT	CAP	CAA	GAA	TTT	ΓGA <i>F</i>	AAGA	ACTI	ΓTG	ATA	AAT(CTT	TTGAT	2640
L	S	D	K			N	٧	N	• •	N	L	• • •	D	•	D	K	•	F	D	
AA		STT(GGT	GAA	AG	ATTI	TAGG	TAT	CAA	TCC	CAGA	ATO	GA7	TTT	CAAA	AAG	ΓTG/	AAAA	CCTT	2760
K	G	S	٧	K	D	L		I		•	Ε	W	I	_	K	•	Ε	N	L	
GC			CGA	CCT	TG	AAAA	TTC	CGT	TAA	AGA	TGT	GAT	CAT	CAA	TCA	AAA	\GGT	ΓΑΑΟ	GGAT	2880
A	• •	S	D.	L	E		•	٧		D	•	I	I	N	Q	K	٧	T	D	
61	AGA	IGCA	AGC	GTT	_		TCT			TTT	CTC	AAA	GGA	IGCA	TTA	rgg(CCCA	ACA	AGCT	3000
V C C	E Tot	Q	A	L	A	D	L	K	N	F	S	K	Ε	Q	L	Α	Q	Q	Α	
ษษ c	161	GAA	166	AAC	CCT	FAGT	CGG	TAA	TGG	GTT	ATC	TCA	AGC	AGA	AGC	CAC	CAAC	TCT	TTCT	3120
U AA	V V V		- G - T C C	1	L	٧	G	N	G	L	S	Q	Α	E	Α	T	T	Ł	S	
HA	LAA	IAA	166	ACI	CAP	AAA	CGA	ACC	CAT	TTA									AGGG	3240
_						N	E	P	I	<u>Y</u>	<u>A</u>	K	٧	N	K	K	K	Α	G	
H I T	אטו מ	ししい	ALI	LAA	ICA	IAAI	AGC.	AAG	IGG	TTT	GGG	TGT	TGT	AGG	GCA	AGC	AGC	GGG	CTTC	3360
CV T	ህ ለተተ	ררר. א	L		Q AAT	I	A	S	G	L	G	٧	٧	G	Q	A	Α	G	F	
UHI E	411	טט <i>ע</i> ^	ICA	UAA V	AAI	IGA	CAA	ICH	CAA	ICA	AGC	GGT	ATC	AGA	AGC	TAA	AGC	AGG	TTT	3480
_ Λ Λ ⁻	L Γርፕ	H ATC	.C.T.	K	1	D	N	L	N	Q	A	V 	S	E	A	K	A	G	F	
HA:	161	אוטו A וטו	וטנ	I GA.	AAG	166	AAAI	AAA	AGI	ACC	TGC	TAG	TTT	GTC	AGC	GAA	ACT	AGA	CAAT	3600
۷ ۸ <i>۱</i>	L L		V.	E Cat) ((**	A	K	K	V	P	A	S	L	S	Α	K	L	D	N	
111/	100	ひみしし	טטע	LAI	וטט	AAC	6CA/	AAA	AAA)		IGA(GTG	GCT	CAA	GCT	CGT	GAA	TGA	TAAG	3720

FIG.4D

HIR N ATAGTTGCGCATAATGTAGGAAGCGTTCCTTTGTCAGAGTATGATAAAATTGGCTTC GTAAAAGACACTAATTCTGGCTTTACGCAATTTTTAACCAATGCATTTTCTACAGCA 1103 V GGTTTCCAAAAATCTTAAAGGATTAAGGAATACCAAAAACGCAAAAACCACCCCTTG 1143 G F Q K S TGAATGCTACCAATTCATGGTATCATATCCCCATACATTCGTATCTAGCGTAGGAAG AACTCTGTAAAATCCCTATTATAGGGACACAGAGTGAGAACCAAACTCTCCCTACGG GACAGACACTAACGAAAGGCTTTGTTCTTTAAAGTCTGCATGGATATTTCCTACCCC GAAAAATCAGAAAAACCATAGGAATTATCACACCTTATAATGCCCAAAAAAAGACGCT ATGCCTTTCAAGGTGAAGAGGCAGATATTATTATTATTCCACCGTGAAAACTTGTG ATCTCATTTTTGTGGGTAAAAAGTCTTTCTTTGAGAATTTATGAAGCGATGAGAAGA CATTCTTCGCTTCAAAACGCTTTCATAAATCTCTCTAAAGCGCTTTATAATCAACAC TTATTAGCGTTACAATTTGAGCCATTCTTTAGCTTGTTTTTCTAGCCAGATCACATC CTGCAAATATCCTACAATAGCATCGCCCGAATGGATGAGTAGGGGGGGTGTTGAAAG TAAAATAATCACTTCGGGAAAATCTTTAAGGGAGTGAAATAATAACGCATGCAAGTT TGCGAAACATTCAAATAGCCTTGTTGTTTCAGGGCATTGTCATAAGCGTTGGATTGG GCTAAAATGCTTGGCTCAATCACGCCCACAATAGGGATTTTGGAATGCTTTTGCATC TTGAAAAAATCCAAAGCCTCTAAGCCAAATTGCTTGATCGTAGTGGGGTCTTTAGTG AGGCTTTTTAAAACGCTAAACCCTCCCACACCGCTATCAAAAACGCCTATTTTCATG TCTTCATTGTCCTTAGTTTGTTGCATTTTAGAATAGACAAAGCTT 5925

FIG. 4E

	K	•	4	1	b	۱۷i	L.	•	1	(1	1		N	P		F	W		L	K		L	٧		N	D	K	,	
AA	CC	AG	AA	GAA	TAT	GA	AAG	AT	TA'	TT(CTO	GAT	TC	GT	TC	AA	GT	TT	TC	CA	CC	AA	GT	TG.	AA	CAA	TG	СТ	3840
N	Q		<	N	M	K			Υ	S	_)		F		K			S	T		K	L		N	N	A		
TC	TT	AT	FA(CTG	CTI	GG	CGA	GA	GA	AAA	AT(GC6	GGA	GC	AT	GG	AA	TC	AA	GA	AC	GT	TA.	AT.	AC.	AAA	AG	GT	3960
S	Y	`	1	C	L	A	R		E	N		1	_	Н		G	I		K	N		٧	N		T	K	G		
<u>CT</u>	AA	A <u>A(</u>	<u> </u>	GAG	GGG	IT	TTT	TA	AT/	ACT	rc(TT	AG	CA	GA	AA	TC	CC	AA	TC	GT	СТ	TT	AG	TA	TTI	GG	GA	4080
																						٠							
TG	TG	CAA	AA(STT	ACG	CC.	ITT	GG	AG/	ATA	AT6	TAG	GT	GT(GA	GA	CC	TG	TA	GG	GA	AT	GC	GT	TG	GAG	iCT	CA	4200
				4GC																									4320
				ГТА																									4440
AC	AA	AGA	T(CAA	GTT	CA	AAA	AA'	TCA	ATA	\GA	\GC	TT	TT	4G	AG	CA	AA'	TT	GA [*]	TC	GC	GC ⁻	TC:	TT	AAC	CA	AA	4560
				SAA																									4680
				CT.																									4800
AT	AT(CTT	T	(GC	GCT	ATT	TTT	GC	AAG	STO	Te	TA	GA	TAG	GG	TA	AT	CT	TT	TC	CA	AA	GA ⁻	ΓΑ	AT(CAT	TA	GA	4920
AA	TA(T	ATA	4GT	GT(SAG	CT	ATA	GC	CC	CT	TT	TTO	GG	GA	AT	TG	AG	TŢ	AT	TT	TG	AC	TT	ΓΑΑ	ΑT	TT	5040
GC	CG(CTC	GC	TAC	GAA	ATT	CC	AC	m	TAG	GG	AA	TG	CGT	rg:	TG	CA	TT	TT	TT	TT	AA	GG(GC(GT/	ATT	TT	TG	5160
GG	CAA	AAA	Te	CT(CCA	TAA	AA	TA(GCC	CT	CA	AT	TT	ITI	ΓG	AG	CG	AT	TA.	AG	GG	AA	AAT	ΓG(CG	TGC	AA	CC	5280
TC	TAA	ICA	TA	TC(GCC	CT(TA.	AA	ATA	CT	TT	CT	TC	AA7	TC/	AA,	AG(GC	AC.	AA	AA	AG	AG/	AA(GT(GGC	TA	AA	5400
				TT																									5520
				GC																									5640
				ACT																									5760
				TA																									5880
																									- '	•			

FIG. 4 F

10 30 50 AAGCTTGCTGTCATGATCACAAAAAACACTAAAAAACATTATTATT<u>AAGGA</u>TACAAAATG M70 90 110 GCAAAAGAAATCAAATTTTCAGATAGTGCGAGAAACCTTTTATTTGAAGGCGTGAGGCAA DSARNL 130 150 170 CTCCATGACGCTGTCAAAGTAACCATGGGGCCAAGAGGCAGGAATGTATTGATCCAAAAA G P R G R TM 210 190 230 AGCTATGGCGCTCCAAGCATCACCAAAGACGCGTGAGCGTGGCTAAAGAGATTGAATTA K D G V S 250 270 290 AGTTGCCCAGTAGCTAACATGGGCGCTCAACTCGTTAAAGAAGTAGCGAGCAAAACCGCT Q L V G A V A S K 310 330 350 GATGCTGCCGGCGATGGCACGACCACAGCGACCGTGCTAGCTTATAGCATTTTTAAAGAA 370 410 390 GGTTTGAGGAATATCACGGCTGGGGCTAACCCTATTGAAGTGAAACGAGGCATGGATAAA A N P450 GCTGCTGAAGCGATCATTAATGAGCTTAAAAAAAGCGAGCAAAAAAGTAGGCGGTAAAGAA K K A S K K V G G K E 490 510 530 GAAATCACCCAAGTGGCGACCATTTCTGCAAACTCCGATCACAATATCGGGAAACTCATC A N S D IS Н 550 570 590 GCTGACGCTATGGAAAAAGTGGGTAAAGACGGCGTGATCACCGTTGAGGAAGCTAAGGGC ADAMEKVGKDGVI TVEEAKG 610 630 650 ATTGAAGATGAATTGGATGTCGTAGAAGGCATGCAATTTGATAGAGGCTACCTCCCCT IEDELDVVEGMQF RGYLSP D

FIG. 5A

		67	70						69	0						710			
TA	Ш	TGT	AAC	GAA	CGC	TGA	GAA	AAT	GAC	CGC	TCA	ATT	GGA	TAA	TGC	TTA	CAT	CCT	ATTT
Υ	F	٧	T	N	Α	Ε	K	M	T	Α	Q	L	D	N	Α	Υ	I	L	L
		73	30	•					75	0						770			
AC	GGA'	TAA	AAA	AAT	СТС	TAG	CAT	GAA	AGA	CAT	TCT	CCC	GCT	ACT	AGA	AAA	AAĊ	CAT	GAAA
T	D	K	K	I	S	S	M	Κ	D	I	L	Р	L	L	Ε	K	T	M	K
								•								•			
		79	90						81	0				Н	ind	III			
GA	GGG	CAA	ACC	GCT	TTT	AAT	CAT	CGC	TGA	AGA	CAT	TGA	GGG	CG <u>A</u>	<u>AGC</u>	<u>TT</u> T	AAC	GAC	TCTA
Ε	G	K	P	L	L	I	I	Α	Έ	D	Ī	Ε	G	E	Α	L	T	T	L
		8	50						87	0						890			
GT	GGT	GAA ⁻	TAA	ATT	AAG	AGG	CGT	GTT	GAA	TAT	CGC	AGC	GGT	TAA	AGC	TCC	AGG	CTT	TGGG
٧	٧	N	K	L	R	G	٧	L	N	I	Α	Α	٧	K	Α	Р	G	F	G
		9.	10						· 93	0						950			
GA	CAG	AAG	AAA	AGA	AAT	GCT	CAA	AGA	CAT	CGC	TAT	TTT	AAC	CGG	CGG	TCA	AGT	CAT	TAGC
D	R	R	K	E	M	L	K	D	I	Α	I	L	T	G	G	Q	٧	I	S
		97	70						99	0					1	010			
GA	AGA	ATT	GGG	CTT	GAG	TCT	AGA	AAA	CGC	TGA	AGT	GGA	GTT	TTT	AGG	CAA	AGC	TGG	AAGG
E	E	L	G	L	S	L	E	N	Α	E	٧	Ε	F	L	G	K	Α	G	R
		10	30						105	0			•••		1	070			
AT	TGT	GAT	TGA	CAA	AGA	CAA	CAC	CAC	GAT	CGT	AGA	TGG	CAA	AGG	CCA	TAG	CGA	TGA	TGTT
I	٧	I	D	K	D	N	T	T	I	٧	D	G	K	G	H	S	D	D	٧
		10							111	•					_	130			
AA	AGA	CAG	AGT	CGC	GCA	GAT	CAA	AAC	CCA	AAT	TGC	CAAG	TAC	GAC	AAG	CGA	TTA	TGA	CAAA
K	D	R	٧	A	Q	I	K	T	Q	I	A	S	T	T	, S	D	Y.	D	K
		11							117	_					_	190			
GA	AAA.	ATT	GCA	AGA	AAG	ATT	GGC	TAA	ACT	CTC	TGG	iCGG	TGT	GGC	TGT	GAT	TAA	AGT	GGGC
E	K	L	Q	E	R	L	A	K	L	S	G	G	٧	A	٧	I	K	٧	G
		12							123						_	250			
																			CGCG
A	A	S	E	٧	Ε	M _.	K	E	K		D	R	V	D	D	A	Ŀ	S	A
		12							129							310			
AC																TCT	CAT	_	CGCG
T	K	Α	Α	٧	E	E	G	I	٧	I	G	G	G	Α	Α	Ĺ	I	R	A.

FIG. 5B

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14 / 14

		13	30						135	50					1	370	:		
GC	TCA	AAA	AGT	GCA	TTT	GAA	TTT	GCA	ACGA	TGA	TGA	AAA	AGT	GGG	CTA	TGA	AAT	CAT	CATG
Α	Q	K	٧	H	L	N	L	Н	D	D	Ε	K	٧	G	Y	Ε	I	I	M
		13	90						141	.0					1	430			
CG	CGC	CAT	TAA	AGC	CCC	ATT	AGC	TCA	TAA	CGC	TAT	CAA	CGC	TGG	TTA	TGA	TGG	CGG	TGTG
R	A	Ī	K	Α	Р	L	Α	Q	I	A	I	N	Α	G	Υ	D	G	G	٧
		14	50		-	_		_	147	'O	-	••			1	490			·
GT	CGT			AGT	AGA	AAA	ACA	CGA			TTT	TGG	TTT	TAA		•	•	TGG	CAAG
٧	٧	N	E	٧	Ε	K	Н	E	G	Н	F	G	F	N	A	S	N	G	K
		15	10	•	_	,,	••	_	153	••	•		•	••	• •	550	••	J	•
TA	TGT			GTT	TAA	AGA	AGG	CAT		_	CCC	CTT	ΑΑΑ	AGT				CGC	ТСТА
Υ	γ	D	М	F	K	F	G	1	ī	D	Р.		K	v	F	R	I	Δ	1
	·	15	70	•	••	_	•	-	159	_	•	-	• • • • • • • • • • • • • • • • • • • •	•	1	610	,	,,	_
CA	AAA		_	TTC	GGT	TTC	AAG	ССТ			`AAC	CAC	AGA	AGC	_			TGΔ	AATC
Q	N	A	V	S	۷.	S	S	1	1	1	T	T	F	Δ	T	V	Н	F	ī
_	••	16	30		•			_	165	ິດ	•	•	_	,,	1	670	",	_	•
AA	AGA		_	AGC	GAC	ፐርር	GGC	ΔΑΤ		_	ΤΔΤ	GGG	TGĞ	СΔТ	_	•••	ΤΔΤ	GGG	AGGC
K	E	F	K	A A	T	P	A	М	Р		М	G	G	M	. G	G		ر 1000	G
••	•	169		•	•	•	,,	••	171	_	• •	u	O	11		730		U	U
ΔΤ	GGG			CVT	CT V	ለርር	ՐՐԲ	СТТ		•	ΤΛС	TAT	۸۸۳	CTG	_			TCC	CTTC
M	G	G	M	M	•#	AUC	CCU		001	111	IAU	HAI	MMI	CIU	CII	I-1 A	AAA	166	
1 1	U	17!	• •	11					177	'n					1.	790			
TC.	τΛΛΙ			ררר	TTT	ሮፒለ	۸Λ۸.	тст		-	TCC	ccc	CCT	CCT	_		ΛΑΛ	ררר	CTCG
1 C	ınnı	116			1 1 1	CIA	MMM	101	CII	1 1 1	וטטו	טטטי	וטטו	GCI	טוו	AIA	AAA	LLU	LILU
		18	10						183	'n									
۲T:	TGT			ΔΤΩ	$C \Lambda \Lambda$	CAA	ΔΛΛ	ΔΤΛ			. V V C	CTT							
O I	, 017	144	776	V I O		CUL	אחח	חוע	101	011	MMU								

FIG. 5C

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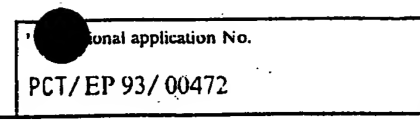
International Application No

PCT/EP 93/00472

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all)6 According to International Patent Classification (IPC) or to both National Classification and IPC Int.Cl. 5 C12N15/31; A61K39/106; C12Q1/68; G01N33/569 C12N15/62; // A61K37/54 **II. FIELDS SEARCHED** Minimum Documentation Searched? Classification System Classification Symbols Int.C1. 5 CO7K ; C12N; C12Q; **G01N** Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched III. DOCUMENTS CONSIDERED TO BE RELEVANT? Citation of Document, 11 with indication, where appropriate, of the relevant passages 12 Category o Relevant to Claim No.13 EP,A,0 367 644 (INSTITUT PASTEUR & INSERM) 1,13-15 9 May 1990 see the whole document 18-23, 26,30-37 & WO,A,9 004 030 cited in the application WO, A, 9 109 049 (RESEARCH EXPLOITATION 1,26, LIMITED) 30-37 27 June 1991 see page 4, line 24 - page 11, line 36 see claims ^o Special categories of cited documents: 10 "T" later document published after the international filing date "A" document defining the general state of the art which is not or priority date and not in conflict with the application but considered to be of particular relevance cited to understand the principle or theory underlying the earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to "L" document which may throw doubts on priority claim(s) or involve an inventive step which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document referring to an oral disclosure, use, exhibition or document is combined with one or more other such docuother means ments, such combination being obvious to a person skilled document published prior to the international filing date but in the art. later than the priority date claimed "&" document member of the same patent family IV. CERTIFICATION Date of the Actual Completion of the International Search Date of Mailing of this International Search Report 18 JUNE 1993 International Searching Authority Signature of Authorized Officer **EUROPEAN PATENT OFFICE** ANDRES S.M. Form PCT/ISA/210 (second short) (January 1925)

III. DOCUME	NTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	
Category o	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.
A	GASTROENTEROLOGY vol. 96, no. 5/2, May 1989, WASHINGTON D.C., US page A101 COVER, T.L. & BLASER, M.J. 'Human serologic response to Campylobacter pylori cytotoxin-associated proteins' see abstract	1-5
A	REVIEWS OF INFECTIOUS DISEASES vol. 13, no. 8, August 1991, pages S686 - S689 LEUNK, R. D. 'PRODUCTION OF A CYTOTOXIN BY HELICOBACTER PYLORI' see the whole document	1-5
P,A	JOURNAŁ OF BIOLOGICAL CHEMISTRY vol. 267, no. 15, 25 May 1992, BALTIMORE, MD US pages 10570 - 10575 COVERT, T.L. & BLASER, M.J. 'Purification and characterization of the vacuolating toxin from Helicobacter pylori' cited in the application see the whole document	1-3





Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	ernational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 23-25 are directed to a method of treatment of (diagnostic method practised on) the human/animal body the search has been carried out and based on the alleged effects of the compound/composition
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Int	ernational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
á. [·	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

EP 9300472 SA 70685

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This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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₩O-A-9109049	27-06-91	None		
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•				
•	e Official Journal of the Euro			